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(54) Title: METHOD FOR THE PREPARATION OF TRANSGENIC PLANTS CHARACTERISED BY GEMINIVIRUS LASTING RESISTANCE

(57) Abstract: The invention relates to a method for the production of transgenic plants durably resistant against *Geminiviruses*, wherein the transgene consists of a suitably modified polynucleotide sequence, derived by the pathogen.





DESCRIPTION

METHOD FOR THE PREPARATION OF TRANSGENIC PLANTS CHARACTERISED BY GEMINIVIRUS LASTING RESISTANCE

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The present invention relates to a method for the preparation of transgenic plants lasting resistant to geminiviruses.

More particularly the invention concerns a method for the preparation of transgenic plants lasting resistant to geminiviruses, wherein the transgene consists of a polynucleotide sequence, derived from the pathogen, suitably modified in order to result in an ineffective target of the post-trascriptional gene silencing induced by geminiviruses.

It is known that geminiviruses are a wide and diversified class of plant viruses that infect several plants of agronomic interest causing serious harvest losses. Such viruses are characterised by virions consisting of two geminate icosahedric particles. Their genome, consisting of one or two circular single-stranded DNA molecules (ssDNA), replicates in the nucleus of infected cells through double stranded intermediates (Hanley-Bowdoin et al., 1999).

The Geminiviridae family is divided in four genera named Mastrevirus, Begomovirus, Curtovirus and Topocuvirus based on the insect vector, the host spectrum and the genome structure (Briddon et al., 1985; Fauquet et al., 2003).

A serious disease of the tomato plant, transmitted by the whitefly *Bemisia tabaci*, is from a long time known as "tomato yellow leaf curl" in the areas of the Middle East, Asian South East and Africa, (Czosnek et al., 1997). This disease, that can cause harvest losses of 100% (Picò et al., 1996; Czosnek et al., 1997), successively spread both throughout the Western Mediterranean, reaching Sardinia, Sicily and Spain (Czosnek et al., 1997), and America (Polston et al., 1997).

Recently the agents of the disease have been identified and isolated, being viruses belonging to the *Geminiviridae* family, genera *Begomovirus*. Phylogenetic studies have highlighted the presence of different viral species related to different geographical origins of the *Begomovirus*: Asia, Africa and America (Czosnek et al., 1997).

The genoma of the Tomato yellow leaf curl Sardinia virus (TYLCSV) species, is monopartite (Kheyr-Pour et al., 1991). The DNA is transcribed

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bidirectionally and contains six open reading frame (ORF), two on the viral strand (V): V1 and V2, and four on the complementary strand (C): C1, C2, C3 and C4, as shown in figure 1. Between the C1 and V2 ORFs there is a non-coding region named intergenic region (IR) analogous to that present in the genome of all *Geminiviridae*. The genomic organization of TYLCSV is structurally similar to that of the bipartite *Begomoviruses* component A such as the tomato golden mosaic virus (TGMV) and the African cassava virus (ACMV). In the case of bipartite *Begomoviruses* the nomenclature of the ORFs present on the component A of the complementary strand is: AL1 or AC1, AL2 or AC2, AL3 or AC3, AL4 or AC4, while on the viral strand AR1 or AV1, AR2 or AV2; on the complementary strand of the component B is: BL1 or BC1 and on the viral strand BR1 or BV1.

Strategies used until now in order to control the infection of the geminiviruses transmitted by the *Bemisia tabaci* are based on the use of expensive fine mesh nets (for the cultivation of fresh-market tomato) and particularly on repeated insecticide treatments (cultivations of both fresh-market and processing tomato). Such strategies result in an increase of the production expenses and represent a serious danger for the health of the agricultural operators and consumer. Furthermore the onset of *Bemisia tabaci* populations resistant to the insecticide imidacloprid has been already reported (Cahill et al., 1996, Williams et al., 1996).

The development of resistant cultivated species represents the most practical and economic way to control viral infections. Classical breeding programs for introducing resistance to geminiviruses that cause the tomato yellow leaf curl were based on the transfer of resistance genes from wild species of *Lycopersicon* to species of cultivated tomato. Thereby lines with variable levels of resistance to TYLCSV have been obtained and commercialized, the best lines showing reduced symptoms and low viral replication. However plants with low and mean levels of resistance represent a potential receptacle for further infections.

Another important aspect to be considered is that the agronomic characteristics of the lines obtained are not always optimal and however reflects those of the genotype of cultivated tomato used in breeding programs.

A tomato line immune to the viruses causing the tomato yellow leaf curl disease, namely, with neither symptoms nor viral DNA replication has not been released yet.

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With the advent of genetic engineering new perspectives were opened up for the introduction of resistance characters against plant viruses. Most strategies are based on the introduction and expression of pathogen-derived sequences in the plant of interest, Pathogen Derived Resistance (PDR) (Sanford & Johnson, 1985; Abel et al., 1986; Tavazza and Lucioli, 1993).

Although such strategies have been successfully applied for the introduction of resistance characters to plant viruses with RNA genome (Beachy, 1997), in the case of geminiviruses, with a DNA genome, the expression of pathogen-derived sequences has produced plants with no lasting resistance and/or tolerance.

The mechanisms that induce virus resistance achieved through the expression of pathogen-derived sequences can be grouped in two wide classes:

a) resistance mediated by the expression of a pathogen protein such as, for instance, the expression of a dominant negative mutant;

b) resistance mediated by the post-transcriptional gene silencing (Baulcombe, 1996; Beachy, 1997; Zaitlin and Palukaitis, 2000).

The post-trascriptional gene silencing is a ubiquitary process in eukaryotes, involving the degradation of specific RNAs following the formation of double strand RNA (dsRNA) molecules having sequences homologous to the target RNA.

Although there may be different contexts able to induce the production of dsRNA homologous to the transgene (transcription of aberrant transgenic RNAs, presence in the transgenic RNA of sufficiently long inverted and repeated sequences, integration of the transgene in the plant genome in inverted and repeated multiple copies), once the dsRNA is produced, the latter is recognised and degraded in short molecules of dsRNA of about 21-26 nucleotides, referred to as siRNA.

The siRNAs are then integrated in a multiprotein complex named RISC, that is able to degrade all RNAs having sequence homology with the siRNAs. The latter ones represent therefore the determining factors of RNA silencing specificity and their presence related to a determined sequence establishes univocally that this RNA sequence is post-transcriptionally silenced.

Therefore, transgenic plants post-transcriptionally silenced for sequences derived from viral RNA genome, are resistant to the

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homologous virus and to viruses with nucleotide sequences closely related to the transgene.

The transgene silencing can be also induced following virus infection.

In fact, viral replication is able to induce silencing of a transgene, initially not silenced, if the nucleotide sequence of the transgene is homologous to a portion of the infecting virus genome. The activation of the silencing mechanism involves the specific degradation of the RNA molecules having sequence homology with the inducer RNA.

As direct consequence, the silencing activation by the virus is associated with a degradation of both transgenic mRNA sequences homologous to the virus and viral genome. This results in the host recovery after an initial infectious step, so that the new vegetative part is proved to be virus free. A peculiar characteristic of the plant tissues that develop subsequently to the recovery phenomenon is that they are highly resistant to a following infection by the same virus.

The resistance mediated by post-transcriptional gene silencing, since based on recognition at the nucleotidic level, confers resistance only against viral isolates closely homologous to the virus genome from which the transgene was derived. Instead, strategies based on the expression of a pathogen protein normally produce plants resistant also to viral strains or isolates not-closely related from a nucleotide point of view.

It is also been shown that the transgene silencing is influenced by the temperature, being inactive at temperatures below 15°C (Szittya et al., 2003). Therefore plants exposed in field conditions at temperature range below 15°C can lose the silencing-mediated resistance.

It must be borne in mind that, although from several years transgenic plants resistant to RNA genome viruses have been achieved through mechanisms based on transgene silencing, so far it is not reported that such strategy can be successfully applied to the geminiviruses (DNA genome-viruses).

It's clear that the best strategy in order to obtain plants resistant to a wide spectrum of geminiviruses is the one in which the interfering product is the protein. It is clear that the width of the resistance spectrum increases the agronomic and commercial value of the produced plant.

Thereby the expression in transgenic plants of dysfunctional variants of geminivirus replicative Rep protein has been used in order to

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obtain plants with greater levels of resistance or immunity against the geminiviruses.

It's known in literature that the expression of a truncated replicative Rep protein (Rep-210) of TYLCSV is able to confer resistance against viral infection, although such resistance is not lasting because the virus is able to overcome it over time.

In tables 1 and 2 are shown the results of the analysis of the resistance of TYLCSV-agroinoculated Rep-210 expressing transgenic plants of *Tomato* 47 x wt (Brunetti et al. 1997) and of *N. benthamiana* line 102.22 (Noris et al. 1996) respectively.

Table 1

	T GDIC 1				
Lycopersicon	Time	N°	infected	%	infected
esculentum	(weeks)	plants/in	oculated	plants/in	oculated
		plants		plants	
Rep-210	4	0/13		0.	
	9	2/13		15	
	18	5/13		38	
Wild-type	4	6/6		100	

Table 2

	1 4510 2				
Nicotiana	Time	N°	infected	%	infected
benthamiana	(weeks)	plants/i	noculated	plants/i	noculated
		plants		plants	
Rep-210	2	4/21		19	
	3	11/21		52	
	4	18/21		86	
Wild-type	2	6/6		100	

From the results reported in tables 1 and 2, it can be clearly inferred that the resistance against TYLCSV mediated by the transgenic expression of a pathogen-derived sequence, is overcome with time.

Similarly, also the resistance induced by the transgenic expression of a dominant negative mutant of Rep of the bipartite geminivirus "African Cassava Mosaic Virus" is overcome with time (Sangaré et al., 1999).

Another example is represented by the transgenic expression of the TYLCV capsid protein in a tomato interspecific hybrid (*Lycopersicon esculentum X L. pennellii*) which confers a partial resistance against viral infection (Kunik et al., 1994). Even in this case the resistance mediated by

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the expression of the capsid protein is not long lasting and it results to be poorly useful from an agronomic point of view.

In the light of the above, is clear the need to have new methods that would allow to use successfully the polynucleotide sequences derived from the geminiviruses in order to obtain long lasting resistant plants against geminiviruses.

The authors of the present invention have now prepared polynucleotide sequences encoding pathogen-derived viral proteins and able to confer virus resistance to the host, suitably modified in order to be ineffective targets of the virus-induced post-transcriptional gene silencing to obtain transgenic plants with lasting levels of resistance against geminiviruses.

In fact during the experiments the authors show that the overcoming of the resistance, and therefore the difficulty to achieve lasting resistance against geminiviruses, is due to the unexpected abilities of the geminiviruses to silence post-transcriptionally the transgene and to spread in a plant in which the transgene, with sequences homologous to the infecting virus, is post-transcriptionally silenced.

As shown in figures 2 and 3, respectively, both in the transgenic plants of *N. benthamiana* line 102.22 and in the plants of *Tomato 47 x wt* the virus ability to overcome the resistance results from the transgene silencing by the same virus and from the unexpected ability of the virus to spread in a silenced plant.

The TYLCSV ability to spread in a plant in which the transgene Rep-210 is post-transcriptionally silenced, is further circumstantiated as set forth in figure 4.

The results show that the transgenic *tomato* plants 47 x 10D (Brunetti et al., 1997), post-transcriptionally silenced before agroinoculation, as shown by the absence of the Rep-210 protein and by the concurrent presence of the transgene-homologous siRNAs, are susceptible to the TYLCSV infection as well as the controls.

From the above it results that, contrary to RNA viruses, the geminivirus is not blocked by an active silencing of viral gene sequences. The above said is not limited to the kind of transgenic plant to be used or the way the virus should be inoculated, through agroinfection or *Bemisia tabaci*. In fact, as shown in table 3, using a reduced number of viruliferous bemisia per plant, so as to infect between 90% and 100% of the control

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plants, about 40% of transgenic plants (line 201) whose transgene is post-trascriptionally silenced, are not or late infected, while at a higher inoculum concentration, all the plants challenged with viruliferous insects are infected similarly to the experiments carried out using agroinoculation.

Table
l able

		abic 0					
Molecular Analysis		Low concentration of		High	concer	ntration	
before inoc	culum	inoculuma			of inoculum ^b		
Transgeni	c plants	2 ^C	3	6	2	3	6
Rep-210	siRNAs	6/15	7/15	8/15	16/21	20/21	21/21
(No)	(Si)	[_			
Not transgenic		2 ^c	3	6	2 .	3	6
Rep-210	siRNAs	11/12	11/12	11/12	8/8	8/8	8/8
(No)	(No)			[{	

a Seven viruliferous insects per plant for 2 days

b Thirty-five viruliferous insects per plant for 5 days

c Weeks after inoculum

Therefore it's important to consider that the viral agroinoculation conditions used for testing the resistance and assessing persistence over time (as shown in figures 2, 3 and 4 and in tables 1 and 2) correspond to high or very high viral pressure conditions. This experimental approach allows to identify transgenic plants with very high resistance levels or immune against the viral infection and therefore of very high commercial value.

Accordingly, the introduction of resistance characters against geminiviruses through the expression of pathogen-derived sequences is limited due to the unexpected ability of the geminiviruses to silence post-trascriptionally the transgene and to spread in the silenced plant.

Furthermore the authors show that the transcripts both of positive (V1 and V2) and negative strand (C1, C2, C3 and C4) of TYLCSV are subjected, during a normal infection on wild-type plants, to the viral post-trascriptional silencing, as shown in figure 5. This results in the impossibility to achieve long-term resistance through expression of sequences derived from the same pathogen, unless these are suitably modified in order not to be a target or to be an ineffective target of the virus-induced post-trascriptional gene silencing. Instead, by introducing in the plant genome a sequence suitably mutated or chosen according to the

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invention it is possible to obtain a long lasting resistance against geminiviruses, unlike that achieved with the known methods.

Therefore it is an object of the present invention a polynucleotide sequence encoding an aminoacidic sequence derived from geminiviruses, said polynucleotide sequence being characterised in that it is not a target or it is an ineffective target of the viral post-trascriptional silencing and having:

- a) a nucleotide homology lower or equal to 90% with respect to the corresponding gene sequence of the geminiviruses against which a resistance is required, preferably lower or equal to 80%, more preferably lower or equal to 70 %;
- b) a continuous homology in the RNA transcript, with respect to the corresponding gene sequence of the geminiviruses against which a resistance is required, lower or equal to 17 nucleotides, preferably lower or equal to 8 nucleotides, more preferably lower or equal to 5 nucleotides;
- c) a maximum length of the sequence containing a single substitution with respect to the corresponding gene sequence of the geminiviruses no longer than 30 nucleotides, preferably no longer than 20 nucleotides, more preferably equal or lower than 9 nucleotides;

said polynucleotide sequence being able to confer to the whole plants, tissues or plant cells therewith transformed, a lasting resistance against the geminiviruses.

The polynucleotide sequences according to the invention can be wild-type or synthetic or produced by mutagenesis and the geminivirus-derived aminoacidic sequences encoded by them are wild-type or mutant sequences that interfere with the viral infection.

Therefore the invention includes polynucleotide sequences of geminivirus either suitably changed or wild-type, such as to differ, at the nucleotidic level, with respect to the corresponding genomic sequence of the geminivirus against which it is required to introduce resistance according to the principles above defined and specified in a), b) and c).

Further object of the present invention is a polynucleotide sequence encoding a geminivirus-derived aminoacidic sequence, said polynucleotide sequence being characterised in that it is not a target or it is an ineffective target of the post-trascriptional silencing and having homology even equal to 100% with respect to the sequence of the geminivirus against which it is required a resistance and being shortened

so as to be underrepresented in the siRNAs population with respect to the original sequence, even if maintaining similar interfering abilities.

The gene sequences from which constructing the polynucleotide sequence according to the invention can derive from the geminiviruses such as, *Mastrevirus*, *Curtovirus*, *Begomovirus*, *Topocuvirus* and particularly can be derived from the species shown in table 4 and their isolates, more particularly from the species of Tomato yellow leaf curl and their isolates shown in table 5.

Table 4

l able 4				
List of geminivirus species	Acronym			
African cassava mosaic virus	ACMV			
Bean calico mosaic virus	BcaMV			
Bean dwarf mosaic virus	BDMV			
Bean golden mosaic virus	BGMV			
Bean golden yellow mosaic virus	BGYMV			
Cabbage leaf curl virus	CaLCuV			
Chilli leaf curl virus	ChiLCuV			
Cotton leaf crumple virus	CLCrV .			
Cotton leaf curl Alabad virus	CLCuAV			
Cotton leaf curl Gezira virus	CLCuGV			
Cotton leaf curl Kokhran virus	CLCuKV			
Cotton leaf curl Multan virus	CLCuMV			
Cotton leaf curl Rajasthan virus	CLCuRV			
Cowpea golden mosaic virus	CPGMV			
Cucurbit leaf curl virus	CuLCuV			
East African cassava mosaic Cameroon virus	EACMCV			
East African cassava mosaic Malawi virus	EACMMV			
EastAfrican cassava mosaic virus	EACMV			
East African cassava mosaic Zanzibar virus	EACMZV			
Indian cassava mosaic virus	ICMV .			
lpomea yellow vein virus	IYVV			
Melon chlorotic leaf curl virus	MCLCuV			
Mungbean yellow mosaic India virus	MYMIV			
Mungbean yellow mosaic virus	MYMV			
Okra yellow vein mosaic virus	OYVMV			
Papaya leaf curl virus	PaLCuV			
Pepper golden mosaic virus	PepGMV			

	
Pepper huasteco yellow vein virus	PHYVV
Pepper leaf curl Bangladesh virus	PepLCBV
Pepper leaf curl virus	PepLCV
Potato yellow mosaic Panama virus	PYMPV
Potato yellow mosaic Trinidad virus	, PYMTV
Potato yellow mosaic virus	PYMV
South African cassava mosaic virus	SACMV
Soybean crinkle leaf virus	SbCLV
Squash leaf curl China virus	SLCCNV
Squash leaf curl virus	SLCV
Squash leaf curl Yunnan virus	SLCYV
Squash mild leaf curl virus	SMLCV
Squash yellow mild mottle virus	SYMMoV
Sri Lankan cassava mosaic virus	SLCMV
Sweet potato leaf curl Georgia virus	SPLCGV
Sweet potato leaf curl virus	SPLCV
Tobacco curly shoot virus	TbCSV
Tobacco leaf curl Japan virus	TbLCJV
Tobacco leaf curl Kochi virus	TbLCKoV
Tobacco leaf curl Yunnan virus	TbLCYNV
Tobacco leaf curl Zimbabwe virus	TbLCZV
Tomato chlorotic mottle virus	ToCMoV
Tomato golden mosaic virus	TGMV
Tomato golden mottle virus	ToGMoV
Tomato leaf curl Bangalore virus	ToLCBV
Tomato leaf curl Bangladesh virus	ToLCBDV
Tomato leaf curl Gujarat virus	ToLCGV
Tomato leaf curl Karnataka virus	ToLCKV
Tomato leaf curl Laos virus	ToLCLV
Tomato leaf curl Malaysia virus	ToLCMV
Tomato leaf curl New Delhi virus	ToLCNDV
Tomato leaf curl Sri Lanka virus	ToLCSLV
Tomato leaf curl Taiwan virus	ToLCTWV
Tomato leaf curl Vietnam virus	ToLCVV
Tomato leaf curl virus	ToLCV
Tomato mosaic Havana virus	ToMHV
Tomato mottle Taino virus	ToMoTV

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Tomato mottle virus	ToMoV
Tomato rugose mosaic virus	ToRMV ·
Tomato severe leaf curl virus	ToSLCV
Tomato severe rugose virus	ToSRV
Tomato yellow leaf curl China virus	TYLCCNV
Tomato yellow leaf curl Gezira virus	TYLCGV
Tomato yellow leaf curl Malaga virus	TYLCMalV
Tomato yellow leaf curl Sardinia virus	TYLCSV
Tomato yellow leaf curl Thailand virus	TYLCTHV
Tomato yellow leaf curl virus	TYLCV
Watermelon chlorotic stunt virus	WmCSV
Wheat dwarf virus	WDV
Maize streak virus	WSV .
Sugarcane streak virus	SSV
Bean yellow dwarf virus	BYDV
Tobacco yellow dwarf virus	TYDV
Tomato pseudo curly top virus	TPCTV
Beet curly top virus	BCTV

Table 5

Species of tomato yellow leaf curl (Fauquet et al., 2003)	Acronym
Tomato yellow leaf curl China virus	TYLCCNV
Tomato yellow leaf curl China virus AF311734	TYLCCNV
Tomato yellow leaf curl China virus - [Y64] AJ457823	TYLCCNV-[Y64]
Tomato yellow leaf curl China virus - Tb [Y10] AJ319675	TYLCCNV-
	. Tp[Y10]
Tomato yellow leaf curl China virus - Tb [Y11] AJ319676	TYLCCNV-
	Tb[Y11]
Tomato yellow leaf curl China virus - To [Y25] AJ457985	TYLCCNV-
	Tb[Y25]
Tomato yellow leaf curl China virus – Tb [Y36] AJ420316	TYLCCNV-
	Tb[Y36]
Tomato yellow leaf curl China virus – Tb [Y38] AJ420317	TYLCCNV-
	Tb[Y38]
Tomato yellow leaf curl China virus - Tb [Y5] AJ319674	TYLCCNV-Tb[Y5]
Tomato yellow leaf curl China virus - Tb [Y8] AJ319677	TYLCÇNV-Tb[Y8]

Tomato yellow leaf curl Gezira virus — [1] AY044137 TYLCGV-[1] Tomato yellow leaf curl Gezira virus — [2] AY044138 TYLCGV-[2] Tomato yellow leaf curl Gezira virus — [2] AY044138 TYLCGV-[2] Tomato yellow leaf curl Gezira virus — [Shambat] AY044139 TYLCGV-[Sha] Tomato yellow leaf curl Malaga virus TYLCMalV Tomato yellow leaf curl Malaga virus AF271234 TYLCMalV Tomato yellow leaf curl Sardinia virus TYLCSV Tomato yellow leaf curl Sardinia virus — Spain [1] Z25751 TYLCSV Tomato yellow leaf curl Sardinia virus — Spain [2] L27708 TYLCSV-ES[2] Tomato yellow leaf curl Sardinia virus — Spain [2] L27708 TYLCSV-ES[2] Tomato yellow leaf curl Sardinia virus — Spain [2] L27708 TYLCSV-Sic Tomato yellow leaf curl Thailand virus — Sicily Z28390 TYLCSV-Sic Tomato yellow leaf curl Thailand virus — [1] X63015, X63016 TYLCTHV Tomato yellow leaf curl Thailand virus — [2] AF141922, AF141897 Tomato yellow leaf curl Thailand virus — [Myanmar] AF206674 TYLCTHV-[1] Tomato yellow leaf curl Thailand virus — [Y72] AJ495812 TYLCTHV-[Y72] Tomato yellow leaf curl virus X15656 TYLCV Tomato yellow leaf curl virus — [Almeria] AJ489258 TYLCV-[Alm] Tomato yellow leaf curl virus — [Almeria] AJ489258 TYLCV-[Alm] Tomato yellow leaf curl virus — [Cuba] AJ223505 TYLCV-[CU] Tomato yellow leaf curl virus — [Dominican Republic] AF024715 TYLCV-[DO] Tomato yellow leaf curl virus — [Saudi Arabia] TYLCV-[SA] Tomato yellow leaf curl virus — [Saudi Arabia] TYLCV-[Shi] Tomato yellow leaf curl virus — [Spain7297] AF071228 TYLCV-[Mid		
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	Tomato yellow leaf curl virus - Mild X76319	TYLCV-MId

Preferably the species of *Begomovirus* are TYLCCNV, TYLCGV, TYLCMalV, TYLCSV, TYLCTHV, TYLCV, ACMV, BGMV, CalCuV, ToCMoV, TGMV, ToGMoV, ToMHV, ToMoTV, ToMoV, ToRMV, ToSLCV, ToSRV, Cotton leaf curl (CLCrV, CLCuAV, ClCuGV, CLCuKV, CLCuMV, CLCuRV), East African cassava mosaic (EACMCV, EACMMV, EACMV, EACMZV), Potato yellow mosaic (PYMPV, PYMTV, PYMV), Squash leaf curl (SLCCNV, SLCV, SLCYV), Sweet potato leaf curl (SPLCGV, SPLCV), Tobacco leaf curl (TbLCJV, TbLCKoV, TbLCZV), Tomato leaf

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curl (ToLCBV, ToLCBDV, ToLCGV, ToLCKV, ToLCLV, ToLCMV, ToLCNDV, ToLCSLV, ToLCTWV, ToLCVV, ToLCV) and isolates thereof.

Other species of preferred geminivirus, belonging to the other genera *Mastrevirus, Curtovirus, Topocuviruses*, are WDV, MSV, SSV, BYDV, TYDV, BCTV and their isolates.

The gene sequence belonging to the genome of the geminiviruses can be the sequence C1/AL1/AC1, C2/AL2/AC2, C3/AL3/AC3, C4/AL4/AC4, V1/AR1/AV1, V2/AR2/AV2, BC1/BL1 and BV1/BR1, particularly, the sequence C1/AL1/AC1 of the previously described geminiviruses and their isolates.

The aminoacidic sequence encoded by the polynucleotide sequence object of the present invention is a pathogen-derived protein able to confer resistance against the geminiviruses to the plants expressing it. Said interfering protein since, according to the invention, is stably expressed, confers a lasting resistance independently from the molecular mechanism by which the protein product is able to induce resistance.

The pathogen-derived protein can be a capsid protein, replication-associated viral protein (Rep), proteins encoded by the genes C2/AL2/AC2, C3/AL3/AC3, C4/AL4/AC4, V2/AR2/AV2, BC1/BL1 and BV1/BR1.

An example of a possible polynucleotide sequence satisfying the above reported requirement is set forth in figures 16A and 16B that show the alignment between the wild-type nucleotide sequence encoding the Rep-210 protein of the TYLCSV and the synthetic nucleotide sequence modified so as not to be a target of the post-trascriptional degradation induced by the infecting virus, where both nucleotide sequences encode the same viral protein.

The plants, tissues or plant cells that can be transformed with this polynucleotide sequences can be tomato, pepper, tobacco, sweet potato, cotton, melon, squash, manioc, potato, bean, soybean, mung bean, beet, sugar cane, corn, wheat.

It is a further object of the present invention a construct comprising an heterologous polynucleotide sequence containing in 5'-3' direction:

a) a polynucleotide sequence acting as promoter in said plant or tissue or transformed cells;

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- b) a non-translated polynucleotide sequence positioned at 5' of the encoding region, belonging or not to the intergenic region of geminivirus;
- c) a polynucleotide sequence according to the invention or a fragment or a variant thereof;
- d) a sequence acting as terminator of transcription, positioned at the 3' with respect to said polynucleotide sequence.

A further object of the present invention is an expression vector comprising the previously described construct.

Further it is an object of the present invention a plant, tissue or transgenic plant cells, progeny thereof as well as seeds comprising in their genome a polynucleotide sequence according to the present invention.

Finally, it is an object of the present invention a method for the preparation of transgenic plants, tissues or plant cells thereof long-lasting resistant to the geminiviruses that comprises the following steps:

- a) "identification" or "selection" of a viral gene sequence encoding an aminoacidic sequence able to confer resistance against geminiviruses:
- b) mutagenesis or "choice" of the viral gene sequence so as to make it an ineffective target of the post-trascriptional silencing induced by infecting geminivirus;
- c) insertion of the geminivirus mutated or chosen gene sequence obtained in step b) through a construct as described previously, in the plant, tissue or plant cell thereof.

With reference to step a) of the method according to the present invention, the term "identification" means the experimental recognition of said viral gene sequence able to confer resistance against geminiviruses, while the term "selection" means the recognition of an already available viral gene sequence able to confer a not lasting resistance against geminiviruses. Accordingly, the method according to the present invention provides furthermore the solution to the problem of the loss of resistance against geminiviruses that occurs through the employment of known sequences.

Particularly, the mutagenesis predicted in step b) is carried out maintaining a nucleotide homology, with respect to the corresponding gene sequence of the geminiviruses against which it is required to obtain a resistance, lower or equal to 90%, preferably lower or equal to 80%, more

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preferably lower or equal to 70%, distributed so as the continuous homology in the transcribed RNA with respect to the corresponding sequence of geminiviruses is lower or equal to 17 nucleotides, preferably lower or equal to 8 nucleotides, more preferably lower or equal to 5 nucleotides and the maximum length of the sequence containing a single substitution with respect to the native gene sequence is not more than 30 nucleotides, preferably not more than 20 nucleotides, more preferably lower or equal to 9 nucleotides.

As the aminoacidic sequence encoded by the polynucleotide sequence identified or selected in step a), according to the present invention, it can be a protein having homology of 100% with respect to the viral wild-type protein.

This mutagenesis includes all those mutations on the nucleotide sequence that don't decrease the ability of the protein to confer resistance against geminivirus. Possible mutations are both silent point mutations and those leading to the substitution with aminoacids having similar biochemical characteristics, or deletions and/or insertions and/or substitutions.

Alternatively, the mutagenesis in step b) of the method according to the present invention consists of deletions of the polynucleotide sequence at the extremities so as said sequence, while maintaining similar interfering abilities, is under-represented with respect to the original sequence, in the natural population of the siRNAs produced by the infecting virus.

Alternatively the "choice" in step b) of the method according to the present invention consists in the recognition of geminivirus wild-type sequences that differ at the nucleotidic level from the geminivirus against which it is required resistance so as not to be a target or to be an ineffective target of the post-trascriptional silencing.

Particularly, the mutagenesis action in step b) of the method according to the present invention can consist of deletions of the 3' or 5' region of the viral gene sequence of step a), until it is identified the minimum region of said gene sequence that is under-represented with respect to the sequence encoding a wild-type protein, in the population of the siRNAs and that said truncated protein maintains the ability to confer resistance against geminiviruses.

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Moreover, the viral gene sequence of step a) of the method according to the present invention can be that of TYLCSV C1/AL1/AC1 gene and the aminoacid sequence can be a protein truncated relatively to the viral wild-type protein such as, for instance, Rep-130.

Among various agronomic applications of the synthetic polynucleotide sequences according to the present invention, of particular interest is their use for obtaining tomato plants resistant to TYLCSV.

In this particular embodiment, the transgenic polynucleotide sequence encoding the truncated viral Rep protein (Rep-210) has been modified through a 3 'deletion resulting in an ineffective target of TYLCSV-induced post-trascriptional gene silencing, while maintaining the ability to confer resistance.

In particular, using stringent hybridizations with radioactive RNA probes, it was identified a transcribed region of the TYLCSV genome that is under-represented in the population of viral-origin siRNAs produced during the infection of the TYLCSV in wild-type plants, as shown in figures 6 and 7. This region corresponds to the first 390 nucleotides of the gene encoding the TYLCSV Rep. Its transcript is an ineffective target of virus-induced post-trascriptional gene silencing, it encodes the Rep-130 protein which results stably expressed, resulting in a lasting resistance.

Therefore, in a particular embodiment of the invention, the aminoacid sequence of geminiviruses, such as the TYLCSV, encoded by the polynucleotide sequence according to the invention can be the truncated Rep-130 protein (SEQ ID No 9). In this case the viral gene sequence made an ineffective or non target of the post-trascriptional silencing, is the SEQ ID No 8.

It's a further object of the present invention a method as described above wherein the mutagenesis in step b) consists of silent point mutations of the viral gene sequence of step a) that maintain the ability of the encoded aminoacid sequence, to confer resistance against geminiviruses and to be an ineffective or non target of the post-trascriptional silencing.

Particularly, the viral gene sequence of step a) can be the V1/AR1/AV1 (CP) gene for instance of TYLCSV (SEQ ID No 12), and in a particular embodiment the viral gene sequence made ineffective or non target of the post-trascriptional silencing is the SEQ ID No 6.

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In addition, the viral gene sequence of step a) can be the TYLCSV C1/AL1/AC1 gene, and in this case the viral gene sequence which was made ineffective or non target of the post-trascriptional silencing can be the SEQ ID No 2 or the SEQ ID No 4.

The present invention now will be described by way of illustrating but not limiting way, according to preferred embodiments thereof, with particular reference to the figures of the enclosed drawings, wherein:

figure 1 shows the genome of tomato yellow leaf curl Sardinia specie virus (TYLCSV). DNA is transcribed bidirectionally and it contains six open reading frames partially or totally overlapping (ORF), two on the viral strand (V) V1 and V2 and four on the complementary strand (C) C1, C2, C3 and C4;

figure 2 shows the expression of Rep-210 protein in TYLCSV-agroinoculated transgenic *N. benthamiana* plants (line 102.22). The symbols (-), (+) and NI mean healthy, infected and non-inoculated plants, respectively. Analysis has been carried out before the agroinoculation with TYLCSV (0 wpi) and four and eight weeks after it (respectively 4 and 8 wpi);

figure 3 shows the expression of the Rep-210 protein and of the transgenic mRNAs in tomato plants (line 47 X wt) before the agroinoculation with TYLCSV (0 wpi) and 22 weeks after it (22 wpi). The symbols (+) and (-) mean plants that are, respectively, infected or healthy at the specified time;

figure 4 shows the analysis of expression of Rep-210 protein and of the siRNAs corresponding to the relative transcript in transgenic tomato plants (47 X 10D line) before TYLCSV agroinoculation. The symbols (+) and (-) on the panel mean the presence and absence of the sense and antisense C1 transgene, respectively; wt means control wild-type plant; while the symbols (+), (-) and NI under the panel of the siRNAs mean the presence (+) and absence (-), respectively, of the virus and the non-agroinoculation (NI) as a control;

figure 5 shows the Northern blot of the small RNAs extracted from TYLCSV-infected wild-type tomato plants (samples 1-4) and non-infected as a control (sample C); M means a molecular weight marker;

figure 6 shows the distribution analysis of the small interfering RNAs with respect to the genome of the TYLCSV. On the top, the linear

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map of the TYLCSV genome; transcripts are pointed out by the arrows (V1 and V2 with the same polarity as the viral genome and C1, C2, C3 and C4 with complementary polarity); open boxes from 1 to 9 positioned under the viral genome map represent the nine PCR fragments (each about three hundred nucleotides in length), in which the genome has been divided and of which the ethidium bromide staining and hybridization with the siRNAs extracted by TYLCSV-infected tomato plants are shown. IR represents a tenth PCR fragment, corresponding to the non-transcribed intergenic region of the TYLCSV genome. The numbers under the panels mean the percentage of hybridization signal with respect to each PCR fragment;

figure 7 shows the analysis of the presence of siRNAs in non-agroinoculated (sample C) or TYLCSV-agroinoculated (samples 1 and 2) wild-type tomato plants at four weeks after inoculation; particularly the siRNAs corresponding to the transcript for Rep-210 (probe A) and for Rep-130 (probe B) were analyzed; the columns 100 and 50 on the left panels correspond to 100 and 50 pg, respectively, of an oligonucleotide homologous to both the A and B probes;

figure 8 shows the nucleotide sequence (SEQ ID No 8) encoding Rep-130 (SEQ ID No 9) of the pTOM130 plasmid. In capital, nucleotides not belonging to the TYLCSV but deriving from cloning; the underlined sequences correspond to BamHI and EcoRI restriction sites used for cloning. The start and stop codons are set forth in boldfaced while the mutations introduced for eliminating the C4 protein expression are in italic and boldfaced characters;

figure 9 shows the Southern blot of total nucleic acids extracts from wild-type *N. benthamiana* protoplasts cotransfected with a TYLCSV infectious clone (pTOM6), along with the plasmid expressing the mutated Rep protein indicated above each column;

figure 10 shows the quantitative analysis of the TYLCSV replication in wild-type *N. benthamiana* protoplasts cotransfected with pTOM6 plasmid along with the plasmid expressing the mutated Rep protein;

figure 11 shows the scheme of pTOM130 plasmid used for obtaining Rep-130-expressing transgenic plants. LB and RB mean left-and right-border respectively; pE35S represents the duplicated *Cauliflower Mosaic Virus* 35S promoter; Rep-130 (SEQ ID No 8) is the sequence encoding the Rep-130 protein (SEQ ID No 9); t35S is the *Cauliflower*

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Mosaic Virus 35S terminator; tNOS is the terminator of the gene encoding the nopalin synthase; nptII is the sequence encoding the neomycin phosphotransferase; pNOS is the promoter of the gene for the nopalin synthase; Kan is the gene for the kanamycin resistance;

figure 12 shows the analysis of the expression of Rep-130 protein (SEQ ID No 9) in transgenic *N. benthamiana* plants transformed with pTOM130 (lines 300-309);

figure 13 shows the analysis of the TYLCSV replication in wild-type (wt) and transgenic *N. benthamiana* protoplasts expressing either the Rep-130 protein (SEQ ID No 9) (lines 300, 301, 303) or the Rep-210 protein (102.22).

figure 14 shows the analysis of the expression of Rep-130 protein (SEQ ID No 9) in transgenic *L. esculentum* plants transformed with pTOM130 (lines 402, 403, 406, 411, 413, 416, 417). A protein extract from a transgenic Rep-130 expressing *N. benthamiana* (line 303) was used as positive control;

figure 15 shows the comparison between a *L. esculentum* transgenic plant transformed with pTOM130 (line 406) expressing Rep-130 and a non-transformed wild-type plant;

figure 16 A and B shows two examples of synthetic sequences encoding Rep-210 (SEQ ID No 2, SEQ ID No 4). The alignment between the wild-type nucleotide sequence encoding TYLCSV Rep-210 protein (Seq_cod_Rep210_wild_type, on the top; SEQ ID No 1) and the synthetic nucleotide sequence modified so as to be an ineffective target of the virus-induced post-trascriptional silencing is shown (Seq_cod_Rep210_silencing_minus, in the bottom; SEQ ID No 2, SEQ ID No 4). In the synthetic sequences, the mutated nucleotides with respect to the wild-type sequence are shaded;

figure 17 shows the analysis of transient expression, by agroinfiltration into *N. benthamiana* leaves, of Rep-210 protein encoded by the plasmid wild-type gene pTOM102(C4 -) and by the synthetic gene Rep-210 silencing minus B (SEQ ID No 4), (plasmid pTOM102 Syn);

figure 18 shows a transient assay for the inhibition of viral replication through co-agroinfiltration of an *A. tumefaciens* strain containing the TYLCSV infectious clone along with *A. tumefaciens* strains containing: a) the pTOM102(C4 -) plasmid expressing the wild-type gene for Rep-210 SEQ ID No 1 (Brunetti et al., 2001), lines 1-3; b) the

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pTOM102Syn plasmid expressing the synthetic gene, for Rep-210 (Rep-210 silencing minus B; SEQ ID No 4) lines 4-6; c) the empty cloning plasmid pBIN19 lines 7-9;

figure 19 shows the analysis of the expression of Rep-210 protein in transgenic *N. benthamiana* plants transformed with pTOM102Syn plasmid containing the synthetic gene for Rep-210, Rep-210-silencing minus B, SEQ ID No 4 (lines 506, 508A and 508B). A protein extract from Rep-210-expressing transgenic tomato plant was used as positive control;

figure 20 shows the analysis of the expression of Rep-210 protein in transgenic *N. benthamiana* plants transformed with pTOM 102 (line 102.22, Noris et al., 1996) or with pTOM 102Syn (line 506) after TYLCSV agroinoculation. Analysis has been performed before (0wpi) and five weeks after (5wpi) TYLCSV agroinoculation;

figure 21 shows the analysis of the infection by "dot-blot" assay at 2, 3, 4, 5 wpi (where wpi means the number of weeks after the agroinoculation) on *N. benthamiana* wild-type (WT) or transgenic plants transformed with either pTOM 102 (line 102.22, Noris et al., 1996) or pTOM 102Syn (line 506);

figure 22 shows an example of synthetic sequence encoding CP. The alignment between the wild-type nucleotide sequence encoding TYLCSV CP (TYLCSV CP, on the top; SEQ ID No 12) and the synthetic nucleotide sequence modified so as to be an extremely ineffective or non target of the virus-induced post-trascriptional degradation (TYLCSV CP silencing minus, in the bottom; SEQ ID No 6) is shown. In the synthetic sequence, the mutated nucleotides with respect to the wild-type sequence are shaded.

<u>EXAMPLE 1</u>: Identification of regions of the TYLCSV genome under-represented in the siRNAs population.

In a natural infection by TYLCSV of wild-type plants, the viral sequences transcribed by both strands of TYLCSV genome are target of post-trascriptional gene silencing as pointed out by the presence of siRNAs homologous to different portions of the genome (figure 5). In figure 5 is shown the Northern blot of total RNAs extracted from the tomato wild-type plants infected by TYLCSV (samples 1-4) and non-infected control (sample C). Probe and the restriction sites used are indicated aside each panel. Also the estimated sizes of siRNAs are set forth.

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In order to evaluate if some regions of the TYLCSV genome constitute a target of post-trascriptional gene silencing less effective than others, it was performed a systematic study of the siRNA distribution with respect to their position on the viral genome. Therefore the TYLCSV genome has been divided in nine contiguous fragments, each of about three hundred base pairs (as drawn in figure 6), obtained by PCR with specific oligonucleotides. The same amount of such fragments has been transferred on nylon filter after agarose gel electrophoresis. Quantification of PCR fragments loaded on agarose gel has been performed by software Aida. The siRNAs produced by a TYLCSV-infected tomato plant have been purified starting from the total RNAs, terminally labelled and used as probe (Szittya et al., 2002) on the filter containing several regions of TYLCSV genome. The different intensity of the hybridization signals, referred to a same amount of loaded fragment, has been assessed through the TYPHOON apparatus (Amersham-Pharmacia). So a different distribution of the siRNAs with respect to the several regions of the viral genome has been detected (fig.6).

<u>EXAMPLE 2</u>: Identification of a region of TYLCSV C1 gene under-represented in the si RNAs population.

In order to identify a region of the TYLCSV C1 gene underrepresented in the siRNAs population, total RNAs (Brunetti et al., 1997) both from healthy and TYLCSV-infected tomato plants have been extracted.

Thirty micrograms of such RNAs have been submitted to 8% denaturing polyacrylamide gel electrophoresis and transferred by capillarity on nylon filter through Northern blot. Two identical replicas have been produced and for each it has been carried out an hybridization with probes corresponding to different portions of the 5' region of the C1 gene, as shown in figure 7. One filter has been hybridized with a probe derived from the 5' portion of C1 gene comprising 42 nt of non-translated leader sequence and the first 630 nucleotides of C1 gene (about 3/5 of C1 gene) (probe A) and the other filter with a probe derived from the 5' portion of C1 gene comprising 42 nt of non-translated leader sequence and the first 390 nucleotides of C1 gene (probe B).

In order to quantitatively compare the results obtained by the two different probes (deriving from two independent labelling), scalar amounts of a same 40mer oligonucleotide complementary to both probes

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have been loaded on both replicas. Columns 100 and 50 correspond to 100 and 50 picograms of such oligonucleotide, respectively. The panels showing the oligonucleotide migration have been set close to the respective panels containing the siRNAs but their position in the figure doesn't correspond to the position on gel, because the oligonucleotide and the siRNAs have different molecular weights.

Both probes after *in vitro* transcription have been submitted to alkaline hydrolysis (Cox et al., 1984) in order to obtain from them fragments with an average length of 75 nucleotides.

The hybridizations have been performed for 16 hours at 39°C in the buffer described by Dalmay et al., 2000. After hybridization the filters have been washed in 2X SSC, 0,2% SDS twice for 10 minutes at 40°C, twice for 10 minutes at 45°C and once for 10 minutes at 50°C.

It is remarkable how the proximal 5' region of the C1 gene in the siRNAs population is under-represented. Particularly, the quantitative analysis of the results performed through the TYPHOON apparatus (Amersham-Pharmacia) revealed that the siRNAs corresponding to this 5' region are about 25% (probe B) with respect to those corresponding to the region extended up to nucleotides encoding the 210 aminoacid (probe A). Said 5' region constitutes therefore an ineffective target for the virus-induced post-trascriptional gene silencing.

These results have been confirmed using the method described in example 1, i.e., where PCR fragments corresponding to the two different regions of the C1 gene were hybridised with the population of siRNA extracted from tomato plants infected by TYLCSV.

EXAMPLE 3: Construction of a polynucleotide sequence of the C1 gene 5' portion encoding the truncated Rep.

As previously pointed out (Brunetti et al., 1997), the Rep-210 transgenic plants show a not long lasting resistance and an altered phenotype.

As can be noticed in figure 1, the C4 gene is nested in the truncated C1 gene in a different reading frame.

It is shown that the transgenic expression of geminivirus C4 gene induces phenotype alterations (Krake et al., 1998).

Therefore, it has been designed several truncated C1 constructs unable to express C4 ORF.

In order to obtain C4 (-) mutants, a stop codon has been introduced in the C4 sequence through the introduction of two point mutations. Particularly, referring to the pTOM130 sequence set forth in figure 8 (SEQ ID No 8), the mutation at nucleotide 233 consists of a trasversion from C to G that converts the TCA codon (encoding serine) of the reading frame encoding C4 in TGA (opal). In addition, the mutation at nucleotide 231 consists of a transition from C to T that restores in the reading frame encoding C1 a leucine codon (CTC becomes TTG, more represented in plant).

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Thereby the translation of the C4 protein is interrupted after only 10 aminoacids, while the aminoacid sequence of the C1 protein remains unchanged. The two introduced mutations have been chosen among many possible mutations based on the criterion to generate a "strong" stop codon in the C4 reading frame, maintaining in the C1 reading frame a leucine codon compatible with codon usage in plants.

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Mutagenesis has been performed by PCR with the following mutated oligonucleotides:

C4 plus.primer (SEQ ID No 10): 5'-CT CAT CTC CAT ATT <u>TTG</u> ATC CAA TTC GAA G-3'

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C4 minus.primer (SEQ ID No 11): 5'-C TTC GAA TTG GAT CAA AAT ATG GAG ATG AG-3 ' (2419-2448 in TYLCV - Kheyr-Pour et al., 1991)

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Each of the two mutated primers has been used along with an external primer in two separate PCR reactions using pGEM102 as template (Brunetti et al., 2001).

Particularly, the external oligonucleotides are Rev and Univ (M13/pUC sequencing primer n.1233 and 1224). From the reaction performed with Univ/C4plus it has been obtained a 537 bp fragment, while from the reaction with Rev/C4minus a 351 bp fragment.

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The obtained products have been used as templates for a following amplification reaction carried out using two external primers.

The obtained PCR product has been digested with EcoRI and BamHI restriction enzymes and cloned into the corresponding sites of pJIT60, thus obtaining pJITR210. In both cases it has been carried out the sequencing to verify clones.

EXAMPLE 4: Identification of the minimal 5' region of TYLCSV C1 gene that when expressed in plant cells is able to inhibit viral replication.

In order to define the minimal 5' terminal region of C1 gene able to confer resistance against TYLCSV, a series of 3'-terminal deletion mutants of C1 gene was cloned in pJIT60 expression vector, resulting in a pJTR series.

The viral sequences have been amplified by PCR with Pfu DNA polymerase (Stratagene), using specific primers containing restriction sites at the ends.

The previously described pJITR210 plasmid, which encodes Rep-210, and contains a stop codon for the internal C4 protein, has been used as template. The fragments obtained by amplification reactions have been digested with BamHI and EcoRI enzymes and cloned in the corresponding sites of pJIT60 resulting in the pJTR series.

All final clones have been sequenced in order to confirm the amplification fidelity and vector-insert junctions. The length and the precise positions of every amplified sequence are set forth in table 6.

The ability of each Rep deletion mutant to confer resistance against TYLCSV has been evaluated through cotrasfection assays of *N. benthamiana* wild-type protoplasts with a TYLCSV infectious clone (pTOM6) along with each mutant, and following analysed for the replication level of the viral genome through Southern blot. The obtained results are set forth in figures 9 and 10.

Table 6

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pJTR210a	42 bp UTR + truncated C1 ORF (630 1985-2656 (1)
	nt) containing C4 encoding region
pJTR210	42 bp UTR + truncated C1 ORF (630 1985-2656 (1)
	nt)
pJTR181	42 bp UTR + truncated C1 ORF (543 2072-2656 (1)
İ	nt)
pJTR156	42 bp UTR + truncated C1 ORF (468 2147-2656 (1)
	nt)
pJTR130	42 bp UTR + truncated C1 ORF (390 2225-2656 (1)
	nt)
pJTR120	42 bp UTR + truncated C1 ORF (360 2255-2656 (1)
l	nt)

			
	pJTR80	42 bp UTR + truncated C1 ORF (240 2375-2656 (1)	
		nt)	
	pJTR54	42 bp UTR + truncated C1 ORF (162 2453-2656 (1)	
٠		nt) ·	

(1) nucleotide numbering of the TYLCSV genome are according to Kheyr-Pour et al. 1991.

The protoplast cotransfection, total nucleic acid extraction and Southern analysis have been performed according to already described methods (Brunetti et al. 2001).

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Total nucleic acids extracts from each protoplast sample have been analysed through Southern blot with a digoxigenin-labelled RNA probe corresponding to the sequence encoding Rep-210, and the pGEM-P plasmid used as control. In particular figure 9 represents a Southern blot of total nucleic acids, where scDNA and ssDNA mean supercoiled and single strand DNA of TYLCSV, respectively.

For an accurate quantitative analysis of the effect of the expression of several truncated forms of Rep on the replication of TYLCSV genome, a Southern analysis has been performed with a ³²P-labelled DNA probe corresponding to the region encoding the first 54 N-terminal aminoacids of Rep and the radioactivity level corresponding to each band detected on filter has been evaluated, through analysis with the Istant Imager apparatus (Canberra, Packard).

Each mutated construct has been assayed in duplicate, in three independent experiments and the value set forth in figure 10 represents the average of two or three cotransfections of the three independent experiments.

The level of TYLCSV replication in the cotrasfection experiments performed with pTOM6 along with pGEM-P control plasmid was considered equal to 100%.

Particularly, figure 10 shows a quantitative analysis, the white and black bars of the histograms represents the amounts of supercoiled and single strand DNA, respectively; error bars indicates the mean standard deviation.

As pointed out by observing figures 9 and 10, the first 130 N-terminal aminoacids of the Rep protein are enough to inhibit almost completely viral replication, while the expression of the first 120 N-terminal aminoacids has no influence.

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<u>EXAMPLE 5</u>: Production of N. benthamiana transgenic plants expressing Rep-130.

The analysis of the ability to inhibit TYLCSV replication by the Rep mutants assessed through transient expression in protoplasts, has revealed that the shortest mutant still effective encodes Rep-130 (SEQ ID No 9) as described in the preceding example.

Also it has been previously revealed in other examples that the proximal 5' portion of C1 gene encoding Rep-130 is a less effective target of post-trascriptional gene silencing compared to sequence encoding Rep-210.

Therefore it has been obtained *N. benthamiana* transgenic plants expressing Rep-130. For this purpose, the pTOM130 plasmid represented in figure 11 has been obtained, by cloning Kpnl-BgIII fragment of pJTR130 into the Kpnl-BamHI sites of pBIN19.

N. benthamiana has been transformed with the A. tumefaciens pGV2260 C58 strain containing pTOM130 plasmid and plants resistant to kanamycin have been regenerated as described (Noris et al. 1996).

The primary transformants have been analysed for the presence of transgene by PCR analysis and for the expression of Rep-130 protein through Western blot, as shown in figure 12.

The protein extracts obtained from transgenic (300-309) or wild-type control (wt) plants have been analysed by Western blot using an anti-TYLCSV Rep rabbit polyclonal primary antibody as described (Noris et al., 1996).

EXAMPLE 6: Plant cells stably transformed with the pTOM130 construct and expressing Rep-130 inhibit TYLCSV replication.

In order to early evaluate the resistance conferred by Rep-130, protoplasts isolated from several primary transgenic Rep-130 expressing *N. benthamiana* plants were transfected with a TYLCSV infectious clone (pTOM6).

Transgenic lines have been chosen for their high Rep-130 expression, as revealed by Western blot analysis (figure 12).

The level of TYLCSV replication in such transgenic protoplasts has been compared with that observed in *N. benthamiana* wild-type and in transgenic protoplasts expressing Rep-210 (line 102.22).

Particularly, figure 13 shows the analysis of TYLCSV replication in wild-type (wt) and transgenic *N. benthamiana* protoplasts expressing

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either Rep-130 (SEQ ID No 9) (lines 300, 301, 303), or Rep-210 (line 102.22). Total nucleic acids extracts from several protoplast samples have been analysed by Southern blot using a digoxigenin-labelled RNA probe corresponding to the Rep-210 transcript.

In order to compare the level of TYLCSV replication in the Rep-130 transgenic protoplasts with that observed in wild-type protoplasts, the total nucleic acids extracted from wild-type protoplasts have been also loaded following 1:10 and 1:50 dilutions, as shown in figure 13.

EXAMPLE 7: Production of tomato transgenic plants expressing Rep-130

The analysis of the ability to inhibit TYLCSV replication by Rep mutants, assessed by transient expression in protoplasts, has revealed that the shortest mutant still effective encodes Rep-130, as described in the previous example.

As previously shown, the proximal 5' portion of C1 gene encoding Rep-130 is an ineffective target of post-trascriptional gene silencing compared with the sequence encoding Rep-210.

Therefore it has been carried out the production of transgenic tomato plants (*Lycopersicon esculentum* cv. Moneymaker) expressing Rep-130.

The tomato has been transformed using *A. tumefaciens* pGV2260 C58 strain containing pTOM130 plasmid (figure 11) and kanamycin-resistant plants were regenerated as described (Brunetti et al. 1997).

The primary transformants have been analysed for the presence of the transgene by PCR analysis and for the expression of Rep-130 protein by Western blot (figure 14).

The protein extracts obtained from transgenic (lines 400) or wild-type control (wt) plants have been analysed by Western blot using an anti-TYLCSV Rep polyclonal rabbit primary antibody as described (Noris et al. 1996).

All transgenic tomato plants expressing Rep-130 protein are phenotypically impossible to distinguish from wild-type plants (figure 15).

EXAMPLE 8: Demonstration of long-lasting resistance against TYLCSV in plants transgenic for the pTOM130 construct expressing Rep130

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In order to assess the lasting of the resistance against TYLCSV conferred by Rep-130 expression, the *N. benthamiana* R1 transgenic plants expressing Rep-130 have been agroinoculated with the *A. tumefaciens* LBA4404 strain containing the TYLCSV infectious clone.

As previously reported, the viral delivery through agroinoculation, used to assay the resistance and evaluate stability over time, corresponds to high or very high viral pressure conditions.

Infection of plants has been assessed at weekly intervals by a "tissue printing" assay, using a digoxigenin-labelled probe specific for the coat protein gene.

The results in table 7 show that, unlike the results described in table 2 concerning transgenic plants expressing Rep-210, transgenic *N. benthamiana* plants expressing Rep-130 protein show a long-lasting resistance when agroinoculated with TYLCSV. This can be deduced by comparison of the resistant plants at 2 and 6 weeks following inoculation.

Table 7 Plants -Time -N°. . .. infected % infected plants / plants/inoculated plants | inoculated plants (weeks) Rep-130 2 0/10 0 4 0/10 0 6 0/10 0 Wild-type 2 9/10 90 4 10/10 100

In addition, it was assessed the stability of the resistance against TYLCSV conferred by Rep-130 expression in transgenic R2 tomato plants.

The plants have been agroinoculated with the *A. tumefaciens* C58C1+pCH32 strain containing TYLCSV infectious clone. The agroinoculation conditions used for assaying the resistance and evaluating stability thereof over time, correspond to high or very high viral pressure conditions.

segregating 3

Rep-130

Wild-type

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The infection has been assessed at intervals of one or two weeks through dot-blot assay, using a radioactively labelled probe specific for the coat protein gene.

The results in table 8 point out that tomato transgenic plants expressing Rep-130 show a long lasting resistance when agroinoculated with an TYLCSV infectious clone. This can be deduced from the comparison of the resistant plants at 3 and 12 weeks after inoculation.

	Table 8				
Plants Time		N° infected	% infected plants /		
	(weeks)	plants/inoculated plants	inoculated plants		
Rep-130	3	0/9	0		
	4	0/9	0		
	5	0/9	0		
	6	0/9	0		
	7	0/9	0		
	8	0/9	0		
	10	0/9	0		
	12	0/9	0		

3/3

4/4

Therefore the resistance is associated to the presence of Rep-130 protein (SEQ ID No 9) and to the ability of TYLCSV-inoculated transgenic plant to stably express Rep-130, because the sequence encoding it is an ineffective target of virus-induced post-trascriptional gene silencing and Rep, even if further mutated, maintains its ability to confer virus resistance.

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EXAMPLE 9: Construction of a synthetic polynucleotide sequence, modified in order not to be or to be an extremely ineffective target of the post-trascriptional gene silencing induced by the infecting virus, encoding TYLCSV Rep-210 protein.

In order to achieve the long lasting expression of TYLCSV Rep-210 protein in transgenic plants, it has been produced, employing the method according to the present invention, a synthetic polynucleotide sequence, able to encode for Rep-210 protein, that is not or is an ineffective target of the post-trascriptional gene silencing induced by the infecting virus.

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In addition, the following criterions have been followed:

- the synthetic polynucleotide sequence is not able to encode the C4 protein, having in positions 231 and 233 the same mutations shown in figure 8, as described for Rep-130 (SEQ ID No 8; SEQ ID No 9).
- the introduced mutations are all silent, namely the protein product encoded by the synthetic polynucleotide sequence matches that encoded by the viral wild-type sequence;
- the mutations were introduced according to the frequency of codon usage in the tomato genes; particularly whenever possible the more frequently used codon in tomato was selected;
- the introduced mutations have been all checked to exclude the possible formation of sequences having a particular function, such as for example polyadenilation or splicing signals, also cryptic.

Following the above described criterions, two synthetic sequences encoding Rep-210 have been designed (figure 16 A and B, SEQ ID No 2, SEQ ID No 3, SEQ ID No 4, SEQ ID No 5).

A non-translated leader sequence at the 5' and a stop codon at the 3' have been added to the sequence of the synthetic Rep-210 silencing minus B gene (SEQ ID No 4).

Particularly, the polynucleotide sequence containing in the 5'-3' order the non-translated leader sequence, the synthetic sequence encoding Rep-210 (figure 16 B; SEQ ID No 4, SEQ ID No 5) and the stop codon has been assembled by PCR starting from oligonucleotides (Prodromou and Laurence, 1992; Stemmer et al., 1995), using a thermostable DNA polymerase with "proof reading" correction activity (Pfu DNA Polymerase, Stratagene and/or Pfx DNA Polymerase, Invitrogen).

The synthetic gene has been subsequently cloned in pJIT60 plasmid under the transcriptional control of 35S promoter of the *Cauliflower mosaic virus* (CaMV) and the transcription termination sequences of the CaMV 35S, producing the pJT60Syn. Then the cassette containing in the 5'-3' order: 35S promoter, Rep-210 synthetic gene, 35S terminator, has been removed from pJT60Syn plasmid by restriction with *Kpnl-Bg/ll* and cloned in the *Kpnl-BamHI* sites of the binary plasmid pBIN19 generating pTOM102Syn.

EXAMPLE 10: Evaluation of the inhibition of viral replication by Rep-210 synthetic gene.

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The correct expression of Rep-210 protein, encoded by the synthetic gene, has been checked through agroinfiltration of *N. benthamiana* leaves, with *A. tumefaciens* C58C1/pCH32 transformed with pTOM102Syn. The strain C58C1/pCH32 transformed with pTOM102 (C4 -) has been used as a positive control, while as negative control the strain C58C1/pCH32 transformed with the binary plasmid pBIN19 was used. Western blot analysis (figure 17) shows the expression of Rep-210 protein encoded by the synthetic gene.

In order to assess the ability of the Rep-210 protein, encoded by pTOM102Syn, to inhibit TYLCSV replication, a transient coagroinfiltration assay has been carried out. *N. benthamiana* leaves have been co-agroinfiltrated with *A. tumefaciens* C58C1/pCH32 strain containing the TYLCSV infectious clone (pTOM6) along with the *A. tumefaciens* C58C1/pCH32 strain containing: a) pTOM102Syn plasmid; b) pTOM102 (C4-) plasmid; c) pBIN19 binary plasmid. The TYLCSV replication has been assessed through Southern analysis of the total nucleic acids extracted from the co-agroinfiltrated tissues 72 hours after the infiltration. This analysis has pointed out that Rep-210 protein expressed by the synthetic gene (pTOM102Syn) and by pTOM102(C4 -) wild-type gene, inhibits TYLCSV replication in a similar manner (figure 18).

EXAMPLE 11: Production of transgenic N. benthamiana plants expressing the synthetic gene for Rep-210.

In order to obtain transgenic *N. benthamiana* plants expressing the synthetic gene for the Rep-210, *N. benthamiana* leaf-discs have been transformed using the A. *tumefaciens* LBA 4404 strain containing pTOM 102Syn plasmid and the kanamycin-resistant plants have been regenerated as described (Noris et al. 1996).

The primary trasformants have been analyzed for the expression of Rep-210 protein by Western blot analysis. Four primary trasformants, 506, 508, 517 and 537 lines accumulating intermediate levels of Rep-210 have been selected for further studies. Figure 19 shows western analysis of proteins extracted from the 506, 508A and 508B plants.

EXAMPLE 12: Stability of the resistance in N. benthamiana transgenic plants for Rep-210 synthetic gene.

The authors have previously shown (Noris et al. 1996; Brunetti et al. 1997) that there is a direct correlation between the amounts of Rep-

210 protein produced by the transgenic plants and resistance against TYLCSV. Transgenic plants transformed with the pTOM102 construct accumulating intermediate levels of Rep-210 protein are susceptible to viral infection, like non-transformed plants (Noris et al. 1996 and unpublished data). The low level of Rep-210 protein in these plants is not enough to completely inhibit viral replication, thus allowing the establishment of an early virus-induced post-trascriptional silencing leading to a drastic reduction in Rep-210 protein accumulation which causes lack of resistance.

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In order to assess if Rep-210 protein encoded by the synthetic gene is not or is an ineffective target of virus-induced post-trascriptional gene silencing and therefore to control over time the viral infection, line 102.22 transgenic plants (R3) and line 506 transgenic plants (R0) expressing similar amount of Rep-210 (figure 20, 0 wpi), have been agroinoculated with the TYLCSV and analysed by dot blot at week intervals for the accumulation of the TYLCSV. As expected (Noris et al., 1996) the transgenic R3 line 102.22 plants (figure 21, 5-8) that accumulate intermediate levels of Rep-210 protein are susceptible as the nontransformed plants (figure 21, 1-4). As shown by dot blot analysis (figure 21, 9-12) the transgenic plants for the synthetic construct (R0 line 506) are resistant to viral infection, accumulating only limited amounts of virus. Interestingly and according to virus inability to post-transcriptionally silence effectively the synthetic gene, Rep-210 was still accumulating 5 weeks after inoculum (figure 20, 5 wpi) and inhibiting over time TYLCSV replication (figure 21, 9-12).

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The results described in the examples point out that it is possible to obtain a long lasting resistance against geminiviruses by expressing in plant a transgene consisting of a pathogen-derived polynucleotide sequence, if the latter is suitably selected or modified in order not to be a target or to be an ineffective target of the post-trascriptional gene silencing by the infecting virus.

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EXAMPLE 13: Construction of a synthetic polynucleotide sequence modified in order not to be a target or to be an extremely ineffective target, of the post-trascriptional degradation induced by the infecting virus, encoding the TYLCSV capsid protein.

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As above reported, the transgenic expression of the TYLCV capsid protein in a interspecific tomato hybrid (*Lycopersicon esculentum X*

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L. pennellii) confers a partial resistance against viral infection (Kunik et al., 1994). Also in this case, the resistance mediated by the expression of the capsid protein is not long lasting.

In order to obtain a stable expression of the TYLCSV capsid protein (CP) by transgenic plants, it has been produced a synthetic polynucleotide sequence, able to encode the CP, which results in an ineffective target of virus-induced post-trascriptional gene silencing.

The synthetic polynucleotide sequence has been designed so as to satisfy the requisite not to be or to be an extremely ineffective target of virus-induced post-trascriptional gene silencing employing the method according to the present invention.

In addition, the following criterions have been followed:

- the introduced mutations are all silent, namely the protein product encoded by the synthetic polynucleotide sequence is exactly matching that encoded by the viral wild-type sequence:
- the introduced mutations consider the frequency of codon usage in the tomato genes; particularly whenever possible the codon more frequently used in tomato is selected;
- the introduced mutations have been all checked to exclude the possible formation of sequences having a particular function, such as for example polyadenilation signals or splicing signals, also cryptic.

Following the above described criterions a synthetic sequence encoding CP has been designed (SEQ ID No 12) (figure 22, TYLCSV CP silencing minus, SEQ ID No 6, SEQ ID No 7).

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CLAIMS

- 1. Polynucleotide sequence encoding a geminivirus-derived aminoacid sequence, said polynucleotide sequence being characterised in that is not or is an ineffective target of the viral post-trascriptional silencing and has:
- a) an homology at nucleotidic level below or equal at 90% with respect to the corresponding gene sequence of the geminiviruses against which the resistance is required;
- b) a continuous homology in the transcribed RNA, with respect to the corresponding gene sequence of the geminiviruses, below or equal to 17 nucleotides;
- c) a maximum length of the sequence containing a single substitution with respect to the corresponding gene sequence of the geminiviruses not higher than 30 nucleotides
- said polynucleotide sequence being able to confer to the plants, tissues or plant cells therewith transformed, a lasting resistance against the geminiviruses.
- 2. Sequence according to claim 1, wherein the homology at the nucleotidic level with respect to the corresponding gene sequence of the geminivirus is below or equal to 80%.
- 3. Sequence according to claim 1, wherein the homology at the nucleotidic level with respect to the corresponding gene sequence of the geminivirus is below or equal to 70%.
- 4. Sequence according to each of the preceding claims, wherein the continuous homology in the transcribed RNA with respect to the gene sequence of the geminiviruses is below or equal to 8 nucleotides.
- 5. Sequence according to each of the preceding claims, wherein the continuous homology in the transcribed RNA with respect to the gene sequence of the geminiviruses is below or equal to 5 nucleotides.
- 6. Sequence according to each of the preceding claims, wherein the maximum length of the sequence containing a single substitution with respect to the corresponding gene sequence of the geminiviruses is not more than 20 nucleotides.
- 7. Sequence according to each of the preceding claims, wherein the maximum length of the sequence containing a single substitution with respect to the corresponding gene sequence of the geminiviruses is not more than 9 nucleotides.

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- 8. Sequence according to each of the preceding claims, wherein the polynucleotide sequence has been mutated or it is a wild-type sequence selected from geminivirus so as to differ at the nucleotidic level with respect to the corresponding genomic sequence of the geminivirus against which a resistance is required according to each of the preceding claims.
- 9. Sequence according to each of the preceding claims, wherein the geminiviruses are selected from the group consisting of species of *Mastrevirus*, *Curtovirus*, *Begomovirus* and *Topocuvirus* and isolates thereof.
- 10. Sequence according to claim 9, wherein *Begomoviruses* species are selected from the group consisting of TYLCCNV, TYLCGV, TYLCMalV, TYLCSV, TYLCTHV, TYLCV, ACMV, BGMV, CalCuV, ToCMoV, TGMV, ToGMoV, ToMHV, ToMoTV, ToMoV, ToRMV, ToSLCV, ToSRV, Cotton leaf curl (CLCrV, CLCuAV, ClCuGV, ClCuKV, CLCuMV, CLCuRV), East African cassava mosaic (EACMCV, EACMMV, EACMV, EACMZV), Potato yellow mosaic (PYMPV, PYMTV, PYMV), Squash leaf curl (SLCCNV, SLCV, SLCYV), Sweet potato leaf curl (SPLCGV, SPLCV), Tobacco leaf curl (TbLCJV, TbLCKoV, TbLCYNV, TbLCZV), Tomato leaf curl (ToLCBV, ToLCBDV, ToLCGV, ToLCKV, ToLCLV, ToLCMV, ToLCNDV, ToLCSLV, ToLCTWV, ToLCVV, ToLCV) and isolates thereof.
 - 11. Sequence according to claim 9, wherein the species belonging to the genus *Mastrevirus*, *Curtovirus*, *Topocuvirus* are selected from the group consisting of WDV, MSV, SSV, BYDV, TYDV, BCTV and isolates thereof.
 - 12. Sequence according to claim 1, wherein the gene sequence is selected from the group consisting of C1/AL1/AC1, C2/AL2/AC2, C3/AL3/AC3, C4/AL4/AC4, V1/AR1/AV1, V2/AR2/AV2, BC1/BL1 and BV1/BR1, belonging to the geminiviruses.
 - 13. Sequence according to claim 12, wherein C1/AL1/AC1 gene sequence is from the geminiviruses, as defined according to claims 10 and 11.
 - 14. Sequence according to claim 1, wherein the geminivirus aminoacidic sequence is a pathogen-derived protein able to confer resistance against the geminiviruses to the plants expressing the protein.
 - 15. Sequence according to claim 14, wherein the protein is selected from the group consisting of capsid protein, replication associated

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viral protein (Rep), proteins encoded by the genes C2/AL2/AC2, C3/AL3/AC3, C4/AL4/AC4, V2/AR2/AV2, BC1/BL1 and BV1/BR1.

- 16. Sequence according to claim 1, wherein the plants, tissues or cells thereof belong to the group consisting of tomato, pepper, tobacco, squash, manioc, sweet potato, cotton, melon, potato, soybean, wine, corn, wheat, sugar cane, bean, beet.
- 17. Polynucleotide sequence encoding a geminivirus-derived aminoacid sequence, said polynucleotide sequence being characterised in that is not or is an ineffective target of the viral post-trascriptional silencing and has homology even equal to 100% with respect to the sequence of the geminivirus against which a resistance is required and is suitably shortened to be under represented in the siRNAs population with respect to the original sequence.
- 18. Construct comprising an heterologous polynucleotide sequence containing in the 5'-3' direction:
- a) a polynucleotide sequence acting as promoter in said plant or tissue or transformed cells;
- b) a non translated polynucleotide sequence positioned 5' of the encoding region;
- c) a polynucleotide sequence as defined according to claims from 1 to 17, a fragment or a variant thereof;
- d) a sequence acting as transcription terminator, positioned 3' with respect to said polynucleotide sequence.
- 19. Expression vector comprising the construct as defined according to claim 18.
- 20. Transgenic plant, tissue or plant cells thereof, comprising in their genome a polynucleotide sequence as defined according to claims from 1 to 17.
- 21. Progeny of the plants and plant tissues according to the claim 20.
- 22. Seed comprising in its genome a polynucleotide sequence as defined according to claims from 1 to 17.
- 23. Method for the preparation of transgenic plants, plant tissue or cells thereof long lasting resistant against geminiviruses including the following steps:
- a) identification or selection of a viral gene sequence encoding an aminoacid sequence able to confer resistance against geminiviruses;

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- b) mutagenesis or choice of the viral gene sequence so as to make it an ineffective target of the post-trascriptional silencing induced by infecting geminivirus;
- c) insertion of the geminivirus gene sequence mutated or chosen in the step b) in the plant, plant tissue or cell thereof using a construct as defined according to claim 18.
- 24. Method according to claim 23, wherein by mutagenesis the homology at the nucleotidic level with respect to the gene sequence of the geminivirus against which resistance is required is maintained below or equal to 90 %, distributed in such way that the continuous homology in the transcribed RNA with respect to the corresponding gene sequence of the geminiviruses is below or equal to 17 nucleotides and the maximum length of the sequence containing a single substitution with respect to the corresponding gene sequence of the geminiviruses is not higher than 30 nucleotides.
- 25. Method according to anyone of the claims 23 and 24, wherein by mutagenesis an homology at the nucleotidic level with respect to the gene sequence of geminivirus is maintained below or equal to 80%.
- 26. Method according to anyone of the claims 23 and 24, wherein by mutagenesis an homology at a nucleotidic level with respect to the gene sequence of geminivirus is maintained below or equal to 70%.
- 27. Method according to anyone of the claims from 23 to 26, wherein by mutagenesis a continuous homology at the nucleotidic level with respect to the gene sequence of geminivirus is maintained below or equal to 8 nucleotides.
- 28. Method according to anyone of the claims from 23 to 27, wherein by mutagenesis a continuous homology at the nucleotidic level with respect to the gene sequence of geminivirus is maintained below or equal to 5 nucleotides.
- 29. Method according to anyone of the claims from 23 to 28, wherein the maximum length of the sequence containing a single substitution with respect to the corresponding gene sequence of geminiviruses is no more than 20 nucleotides.
- 30. Method according to anyone of the claims from 23 to 29, wherein the maximum length of the sequence containing a single substitution with respect to the corresponding gene sequence of geminiviruses is no more than 9 nucleotides.

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- 31. Method according to anyone of the claims from 23 to 30, wherein the mutagenesis consists of silent point mutations or deletions and/or insertions and/or substitutions.
- 32. Method according to anyone of the claims from 23 to 31, wherein the mutagenesis in step b) consists of deletions of the 5' or 3' regions of the viral gene sequence of step a) until the identification of the minimum region of said gene sequence that is under represented with respect to the sequence encoding the wild-type protein, in the population of the interfering siRNAs and that said truncated protein maintains the ability to confer resistance against geminiviruses.
- 33. Method according to claim 32, wherein the viral gene sequence of step a) is the C1/AL1/AC1 gene.
- 34. Method according to claim 32, wherein the C1/AL1/AC1 gene is a TYLCSV gene.
- 35. Method according to claim 32, wherein the aminoacid sequence is a truncated protein with respect to the viral wild-type protein.
- 36. Method according to anyone of the claims from 32 to 35 in which the viral gene sequence made not target or ineffective target of the post-trascriptional silencing is the SEQ ID No 8.
- 37. Method according to anyone of the claims from 32 to 36, wherein the truncated protein is Rep-130 (SEQ ID No 9).
- 38. Method according to anyone of the claims from 23 to 31, wherein the mutagenesis in step b) consists of silent point mutations of the viral gene sequence of step a) to maintain the ability of the aminoacid sequence, encoded by the same, to confer resistance against geminiviruses and not to be or to be an ineffective target of the post-trascriptional silencing.
- 39. Method according to claim 38, wherein the viral gene sequence of step a) is the V1/AR1/AV1 (CP) gene.
- 40. Method according to claim 39, wherein the V1/AR1/AV1 (CP) gene is a TYLCSV gene.
- 41. Method according to anyone of the claims from 38 to 40, wherein the viral gene sequence made not target or ineffective target of the post-trascriptional silencing is the SEQ ID No 8.
- 42. Method according to claim 38, wherein the viral gene sequence of step a) is C1/AL1/AC1 of TYLCSV.

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43. Method according to claim 42, wherein the viral gene sequence made not target or ineffective target of the post-trascriptional silencing is the SEQ ID No 2 or the SEQ ID No 4.

ABSTRACT OF THE DISCLOSURE

The invention relates to a method for the production of transgenic plants durably resistant against Geminiviruses, wherein the transgene consists of a suitably modified polynucleotide sequence, derived by the pathogen.

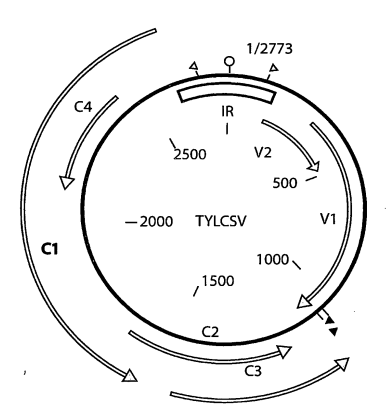


Fig. 1

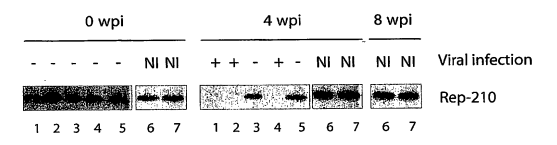
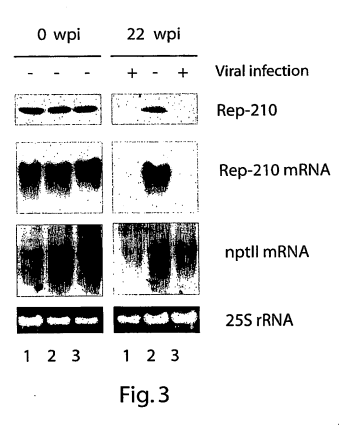


Fig. 2



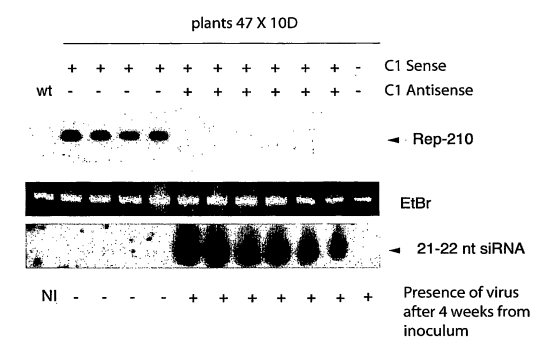


Fig. 4

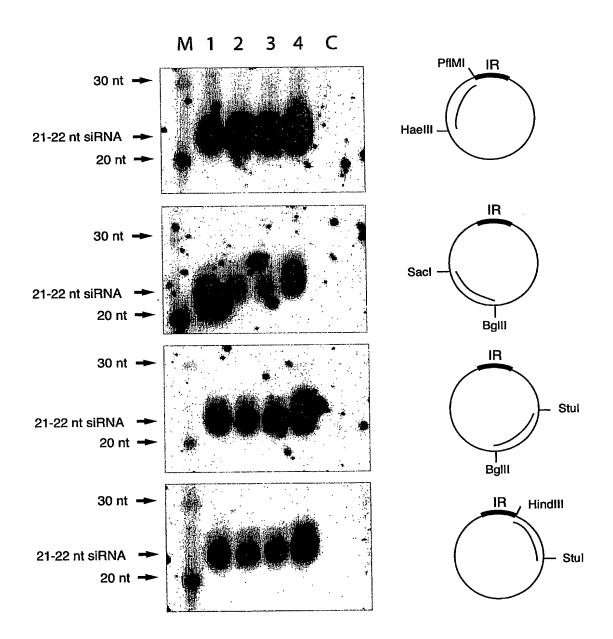


Fig. 5

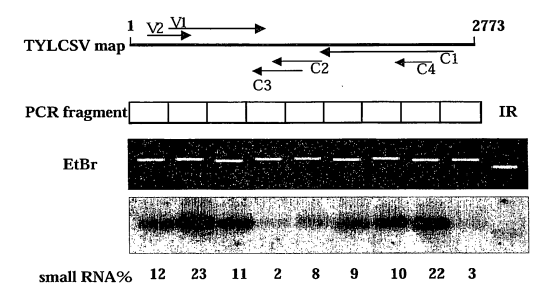


Fig. 6

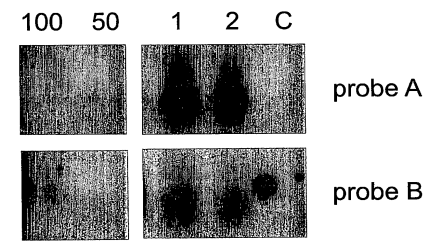


Fig. 7

7/16

SEQ ID No 8

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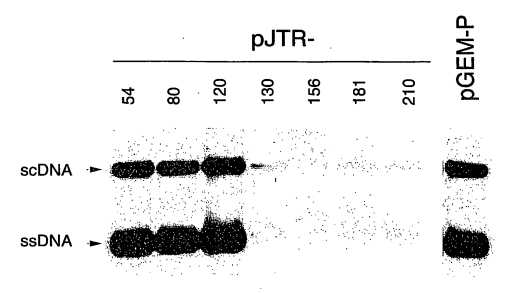


Fig. 9

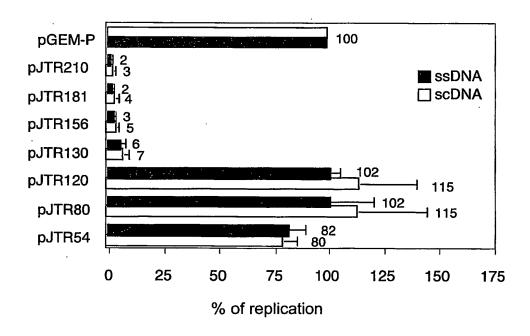


Fig. 10

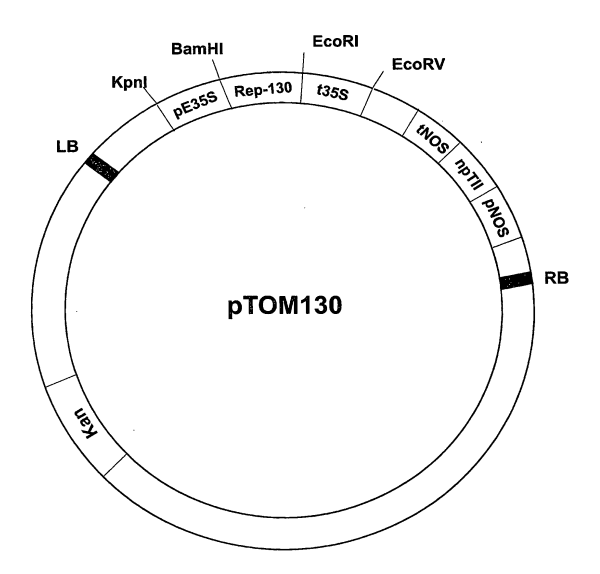


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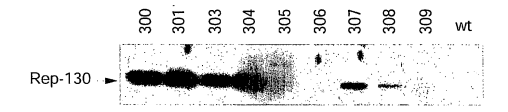


Fig. 12

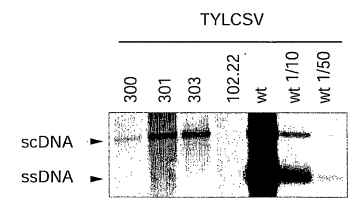


Fig. 13

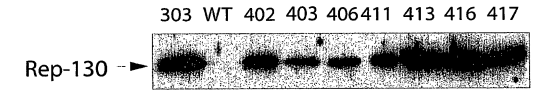


Fig. 14

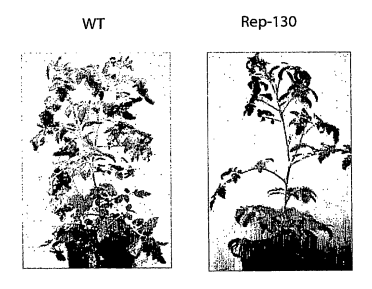


Fig. 15

					12/			_	_	_	
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1 AT G C G A A G A T C A G G T C G T T T A G T A T C A A G G C T A A A A A T T A T C C T T A C A T A T	io 100 100 100 100 100 100 100 100 100 10	140 140 140 140 140 140 140 140 140 140	170 180 180 190 200 200 200 200 200 180 200 200 180 200 180 200 200 180 CCTCATCTCCAATTCCAATTCCAATTACAATTACAATTACAATTACAATTACAATTACAATTACAATTACAATTACAATTACAATTACAATTACAATTAAATTACAATTAAATTACAATTAAATTACAATTAAAA	221 GATTCTTCGACCTGGTATCCCCAACCAGGTCAGCACATTTCCATCCGAACATTCA 221 GTTTCTTGGACCTGGTATCCCCGAACCAGGTCAGCACATTTCATCGAACATTCCA 221 GTTTCTTGGACCTGGTGTCCCCGAACCAGGATCAGCGCATTTCATCGAACATGCA	276 GGGAGCTAAATCGAGCTCCGACGTCAAGTCCTATATCGACAAGGGAGATGTT 276 GGGAGCTAAATCGAGTTCGAGGTCGAGGTGTTATATCGACAAGGACGGAGATGTT 276 GGGTGCTAAGTCGAGTTTCGGAGCTTGAAGTCGATGGACAAGGACGGGATGTG	340 340 340 340 340 340 340 340 340 340	3go 4go 4go 4go 4go 4go 4go 4go 4go 3go 4go 386 C A G G C A G G C A G G C A G G C A G G C A G G C T C T C T C T C T C A G G	490 490 490 490 490 490 490 441 TGATATTCATAATATATATATATATATATATATATATATA	Sport	ssorting to the sport of the s	610

Fig. 16A

				13	3/16					
Rep-210 wild type (SEQ ID No 1) Rep-210 silencing minus B (SEQ ID No 4)	Rep-210 wild type (SEQ ID No 1) Rep-210 silencing minus B(SEQ ID No 4)	Rep-210 wild type (SEQ ID No 1) Rep-210 silencing minus B(SEQ ID No 4)	Rep-210 wild type (SEQ ID No 1) Rep-210 silencing minus B(SEQ ID No 4)	Rep-210 wild type (SEQ ID No 1) Rep-210 silencing minus B(SEQ ID No 4)	Rep-210 wild type (SSQ ID No 1) Rep-210 silencing minus B(SSQ ID No 4)	Rep-210 wild type (SEQ ID No 1) Rep-210 silencing minus B(SEQ ID No 4)	Rep-210 wild type (SEQ ID No 1) Rep-210 silencing minus B(SEQ ID No 4)	Rep-210 wild type (SEQ ID No 1) Rep-210 silencing minus B(SEQ ID No 4)	Rep-210 wild type (SEQ ID No 1) Rep-210 silencing minus B(SEQ ID No 4)	Rep-210 wild type (SEQ ID No 1) Rep-210 silencing minus B(SEQ ID No 4)
30 40 50 60 60 CONDANTATITCCITACATATCCCAAA CAAAAGAATTAATTCATAAAAAAAAAA	90 140 150 170 170 170 170 170 170 170 170 170 17	150 150 160 170 190 ACTACTCCATATT ACTECATGAGAACGTCATCTTCCATATT ACTECATGATGATGATGATGATGATGATGATGATGATGATGATGA	210 220 230 240 ATTGTACCAATCAACGATTCTTCGACCTGGTATCC AEGTGGGACCAAGGCAACGEGTTCGTTGGACCTEGGTATCC	270 280 290 300 GAACATTCAGGGAGCTAAATCGAGCTCCGACGTC MAACATEGCAGGEGGCTAAGTCGAGETTCEGACGTEG	330 340 350 360 360 T CT T CT T C G A C G G A C T T T C C A G A C G G G A C G T A C T T T T C A G A T T A G A C G G T A C T T T T C A G A T A C G G T A C T T T T T A C A C G G T A C T T T T T A C A C G G T A C G G T A C T T T T T A C A T T A C G G T A C G G T A C T T T T T A C A T T A C G G T A C G G T A C G G T A C A C A C A C A C A C A C A C A C A	390 400 420 AGCCAACGCTTACGCAAAGGCAATTAACGCA AGCMAACGAMGCMTACGCMAAGGCMATEAACGCM	450 460 470 480 ANTERAGEOUTOTA ANTERICATE AN	sio 520 530 540 AGATATCCAGGTGCCTCCGGCACCTTAT EGGAEGAAEGTTCCAEGTGCCTCGGCEGTAAT	Sio Sio Garcas Sio Sio Sio Carcas Carcas Sio Sio Carcas Carcas Sio	(で)
1 ATGCCAAGATCAGGTCGTTTAGTAT 1 ATGCCGAAGATCGGGGAAGGAT	10 10 10 10 10 10 10 10 10 10 10 10 10 1	130 140 121 AAATTCATCAAAATTTGCAGAGA 121 AAGGTTGTTCATTAAGGATTGTAGGGA	190 200 181 CTCATCCAATTCGAAGGAAATACAA 181 ETGATACAEGTCGAAEGTAEDAA	250 260 241 CCAACCAGGTCAGCACATTTCCATCC 241 CCMACCAGMTCAGC©CATTMCATCC	310 301 AAGTCCTATATCGACAAGGACGAGA 301 AAGTCATAGATTGACAAAGACGGGGA	370 361 CGATCTGCTAGGGGAGGACACAGAC 361 CGATCGGCTAGAGGAGGAGAACAAACA	430 421 GGAAGTAAGTCGCAGGCTCTTGATGT 421 GGATGGAAGTCAGGCACTTGAEGGT	490 481 CATTTCATAATAAATAGTATTT 481 CATTEGCATAAEAAEGAGEAATTT	541 GTTTCTCCTTTTATCTTCTTCTTTT 541 GTTTCTTCTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	610 FCCGAGAACGTCATGGATGCCGCTGC 601 TCTGATAACGTTAATGGALGGCCGCLAGC

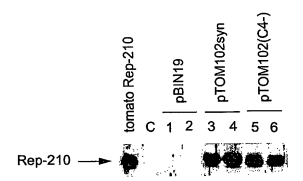


Fig. 17

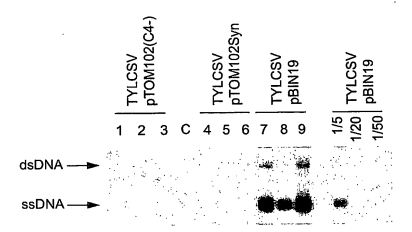


Fig. 18

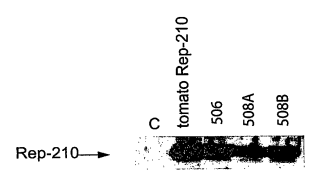


Fig. 19

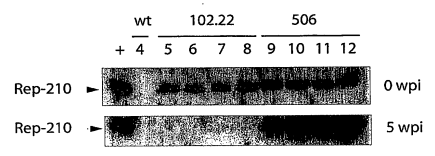
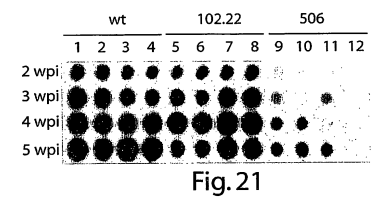


Fig. 20



							16/	16										
	TYLCSV CP (SEQ ID No 12) TYLCSV CP silencing minus (SEQ ID No 6)	TYLCSV CP (SEQ ID No 12)	ON OT Pack course forcements to account	TYICSV CP (SEQ ID No 12) TYICSV CP silencing minus (SEQ ID No 6)	iyicsv cp (seq id no 12)	TYLCSV CP silencing minus (SEQ ID No 6)	TYLCSV CP (SEQ ID No 12) TYLCSV CP silencing minus (SEQ ID No 6)		TYLCSV CP (SEQ ID No 12) TYLCSV CP silencing minus (SEQ ID No 6)	;	TYLCSV CP (SEQ ID No 12) TYLCSV CP silencing minus (SEQ ID No 6)		TYLCSV CP (SEQ ID No 12) TYLCSV CP silencing mirnus (SEQ ID No 6)		TYLCSV CP (SEQ ID No 12) TYLCSV CP silencing mimus (SEQ ID No 6)		TYLCSV CP (SEQ ID No 12) TYLCSV CP silencing minus (SEQ ID No 6)	
J 08 0/	CTGAACTTC	150 150 160 160 160 160 160 160 160 160 160 16	230	GTGCAGTCGTACGAGCAGCGT GTGCANTCGTANGAGCANCGT	300 310 320 320 320 GTAGATTAGAGTTGGTAA	ACTCACAGGGTGG	AAAACAAAATCATACTAAC BAAACAGAATCAAA	460 470 480	G T C A A G T T T T A A C A T G T T T S G G C A G G T G T T T A A A A A T G T T C	05'5	GAGGAAGTTCATGCTACGGT GAGAAAGTT <mark>O</mark> CATGC <u>A</u> ACGGT	630	A C C C A T G T A G T T T A T A A A I A C II C A T G T G T A T A A A I	710	rgtactcatgcttctaaccca rgtactgcatgctataatcca			
9 05	TCGTCGAAG GCGTAGGAG	130 GTCGATCATGGACTTACA		TGTGAAGGTCCCTGTAAA TG C GAAGG <u>A</u> CCCTG G AAG	290 TGTAACTAGGGGTTC <u>T</u> G	TGTGACTAGNGGTTCAG	GGATGGATGAAACATAA GGATGGA G AA TA T A	450 46	ACTAGTCCTATGGATTTTG	530	GATAGGTATCAAGTAAT GATAG <mark>A</mark> TATCA <mark>G</mark> GT G AT	610 620	AAGAGTTTTTAAAATTAATA AABAGATTOTTAAGATOAAG	0690	TTGTTATTGTATATGGCTT		TGTACAATTAA TGTGACAAAOTAG	Fig. 22
30 40	ACTAATTCAACGCCCG	110 120 120 120 120 120 120 120 120 120	190	CCTGATGTACCTCCGGGT	TGTGCGTTGTGTAGTGA		ATATATTAGGAAAGAT ACATATTGGGTAAGAT	430 440	GACCGAAGGCCTTATGGA GAICGAAGACCATACGGA	510 520	GGTGAAGAACGACTTACGG TGTGAABAATGANTTGCG	2590	A G G A G C A G T G T T T G C T G A A G G C A A T G T O T T C T G	089 019	A A T C A T A C T G A G A A T G C C A A T C A C A C T G A A A A T G C II	750	TATITATTTTATGATGCT TATETATTTETATGAEGCT	
10 20	TGCCGAAG TGCCAAG	ACCAGCGTGCTGCCCA	170 180	GATGTACAGAATGTACAG GATGTATAGAATGTATC	ATGACGTCAAGCATACCGGTGT	ACGARGTGAAGCACACACGGAG	CGTTTTTGTATCAAGTCA	410 420	GGT TATGTTTT CCTTG1	490 500	ATAATGAACCCAGTACTGCTAC ATAACGAGCC <mark>ATC</mark> TACTGC <mark>D</mark> AC	570 580	GTTGGAGGTCCGTCAGGGATGA GTTGGTGGTCCTTCTGGGATGA	099 059	CCAAGAGCAGGCGAAGTATGAA CCAGGAGCAAGTATGAG	730 740	TGTACGCTACGTTGAAAATACG TTTACGCAACGCTTAAAGATGCG	
ı	44	⊷ <u>r</u>		161 G	0	241 G	321 A	I	401 A	1 1	481 G 481 G	ŀ	561 561 761	1	641 A 641 A	i	721 721 G	

PCT/IT2004/000287

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gaa Glu	ctt Leu 50	cat His	gag Glu	aat Asn	gga Gly	gaa Glu 55	cca Pro	cat His	ctt Leu	cat His	atc Ile 60	ttg Leu	ata Ile	cag Gln	ttc Phe	1	92
gaa Glu 65	ggc Gly	aag Lys	tat Tyr	aac Asn	tgc cys 70	acc Thr	aac _. Asn	caa Gln	cgt Arg	ttc Phe 75	ttt Phe	gac Asp	ctt Leu	gtg Val	tcc Ser 80	2	40
cct Pro	acc Thr	aga Arg	tca Ser	gcc Ala 85	cat His	ttt Phe	cat His	cca Pro	aac Asn 90	atc Ile	cag Gln	ggt Gly	gct Ala	aag Lys 95	tcg Ser	2	88
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cat His	ttc Phe	cat His	aac Asn	ato 11e 165	Asn	agc Ser	aat Asn	ttg Leu	gac Asp 170	Lys	gtg Val	ttc Phe	caa Gln	gtg Val 175	cca Pro	. !	528
ccg Pro	gct Ala	cct Pro	tac Tyr 180	· val	tca Ser	cct Pro	ttc Phe	tta Leu 185	Ser	tct Ser	tca Ser	ttt Phe	gat Asp 190	GII	gtt Val	;	576
cca Pro	gat Asp	gag Gli 195	ı Let	gag Gli	cat His	tgg Trp	gtç Val 200	Ser	gaa Glu	aac Asr	gtt Val	atg Met 205	. ASL	gco Ala	gca Ala		624
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		3		•													
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Thr Asn Leu Gln Thr Pro Thr Asn Lys Leu Phe Ile Lys Ile Cys Arg 35 40 45 Glu Leu His Glu Asn Gly Glu Pro His Leu His Ile Leu Ile Gln Phe 50 60 Glu Gly Lys Tyr Asn Cys Thr Asn Gln Arg Phe Phe Asp Leu Val Ser Pro Thr Arg Ser Ala His Phe His Pro Asn Ile Gln Gly Ala Lys Ser 85 90 95 Ser Ser Asp Val Lys Ser Tyr Ile Asp Lys Asp Gly Asp Val Leu Glu 100 105Trp Gly Thr Phe Gln Ile Asp Gly Arg Ser Ala Arg Gly Gln Gln 115 120 125 Thr Ala Asn Asp Ala Tyr Ala Lys Ala Ile Asn Ala Gly Ser Lys Ser 130 140 Gln Ala Leu Asp Val Ile Lys Glu Leu Ala Pro Arg Asp Tyr Val Leu 145 - 150 155 160 His Phe His Asn Ile Asn Ser Asn Leu Asp Lys Val Phe Gln Val Pro 175 175 Pro Ala Pro Tyr Val Ser Pro Phe Leu Ser Ser Ser Phe Asp Gln Val 180 185 190 Pro Asp Glu Leu Glu His Trp Val Ser Glu Asn Val Met Asp Ala 195 200 Ala Arg 210 <210> 4 <211> 630 <212> DNA Artificial <223> TYLCSV Rep-210 modified sequence <220> <221> <222> CDS (1)..(630) atg cct aga tcc gga agg ttt agc atc aaa gct aag aat tac ttc ttg Met Pro Arg Ser Gly Arg Phe Ser Ile Lys Ala Lys Asn Tyr Phe Leu 10

Pagina 3

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cct acc aga tca gcc cat ttt cat cca aac atc cag ggt gct aag tcg Pro Thr Arg Ser Ala His Phe His Pro Asn Ile Gln Gly Ala Lys Ser 85 90 95	288
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ccg gct cct tac gtt tca cct ttc tta agt tct tca ttt gat cag gtt Pro Ala Pro Tyr Val Ser Pro Phe Leu Ser Ser Ser Phe Asp Gln Val 180 185 190	576
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20

25

Thr Asn Leu Gln Thr Pro Thr Asn Lys Leu Phe Ile Lys Ile Cys Arg 35 40 45Glu Leu His Glu Asn Gly Glu Pro His Leu His Ile Leu Ile Gln Phe 50 60 Glu Gly Lys Tyr Asn Cys Thr Asn Gln Arg Phe Phe Asp Leu Val Ser 65 70 75 80 Pro Thr Arg Ser Ala His Phe His Pro Asn Ile Gln Gly Ala Lys Ser 85 90 95 Ser Ser Asp Val Lys Ser Tyr Ile Asp Lys Asp Gly Asp Val Leu Glu 100 105Trp Gly Thr Phe Gln Ile Asp Gly Arg Ser Ala Arg Gly Gln Gln 115 125 Thr Ala Asn Asp Ala Tyr Ala Lys Ala Ile Asn Ala Gly Ser Lys Ser 130 135 140 Gln Ala Leu Asp Val Ile Lys Glu Leu Ala Pro Arg Asp Tyr Val Leu 145 150 160 His Phe His Asn Ile Asn Ser Asn Leu Asp Lys Val Phe Gln Val Pro 165 170 175

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tct Ser	att Ile	tac Tyr 115	ata Ile	ttg Leu	ggt Gly	aag Lys	atc Ile 120	tgg Trp	atg Met	gac Asp	gag Glu	aat Asn 125	atc Ile	aag Lys	aaa Lys	384
cag Gln	aat Asn 130	His	act Thr	aat Asn	cag Ģln	gtt Val 135	atg Met	ttc Phe	ttt Phe	ctt Leu	gtg Val 140	cga Arg	gat Asp	cga Arg	aga Arg	432
cca Pro 145	Tyr	gga Gly	acc Thr	agc Ser	cca Pro 150	atg Met	gac Asp	ttc Phe	ggc Gly	cag Gln 155	gtg Val	ttt Phe	aat Asn	atg Met	ttc Phe 160	480
gat Asp	aac Asn	gag Glu	cca Pro	tct Ser 165	act Thr	gca Ala	act Thr	gtg Val	aaa Lys 170	Asn	gat Asp	ttg Leu	cgt Arg	gat Asp 175	aga Arg	528
tat Tyr	cag Gln	gtg Val	atg Met 180	Arg	aag Lys	ttc Phe	cat His	gca Ala 185	Thr	gtg Val	gtt Val	ggt Gly	ggt Gly 190	Pro	tct	576
gga G1y	atg Met	aaa Lys 195	Glu	caa Gln	tgt Cys	ctt Leu	ctg Leu 200	Lys	aga Arg	ttc Phe	ttt Phe	aag Lys 205	716	aac Asn	act Thr	624
cat His	gto Val 210	٧a٦	tat Tyr	aac Asn	cac	cag Gln 215	Glu	caa Gln	gcg Ala	aaa Lys	tat Tyr 220	Glu	aat Asn	cac His	act Thr	672
gaa Glu 225	ı Asr	gct Ala	ttg Lei	ı ttg ı Leu	tta Leu 230	⊢⊤yr	atg Met	gco : Ala	tgt Cys	acc Thr 235	HIS	gca Ala	tct Ser	aat Asr	cca Pro 240	720
gtt Val	tac Tyr	gca Ala	acg Thr	ctt Leu 245	Lys	ato	cgt Arg	ato	tat Tyr 250	' Phe	tat Tyr	gac Asp	gct Ala	gtg Val 255	aca Thr	768
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Pro Pro Gly Cys Glu Gly Pro Cys Lys Val Gln Ser Tyr Glu Gln Arg 65 70 75

Asp Asp Val Lys His Thr Gly Val Val Arg Cys Val Ser Asp Val Thr $85 \hspace{1.5cm} 90 \hspace{1.5cm} 95$

Arg Gly Ser Gly Ile Thr His Arg Val Gly Lys Arg Phe Cys Ile Lys 100 105

Ser Ile Tyr Ile Leu Gly Lys Ile Trp Met Asp Glu Asn Ile Lys Lys 115 . 120 125

Gln Asn His Thr Asn Gln Val Met Phe Phe Leu Val Arg Asp Arg Arg 130 140

Pro Tyr Gly Thr Ser Pro Met Asp Phe Gly Gln Val Phe Asn Met Phe 145 $\,$ 150 $\,$ 155 $\,$ 160

Asp Asn Glu Pro Ser Thr Ala Thr Val Lys Asn Asp Leu Arg Asp Arg 165 170 175

Tyr Gln Val Met Arg Lys Phe His Ala Thr Val Val Gly Gly Pro Ser 180 185 190

Gly Met Lys Glu Gln Cys Leu Leu Lys Arg Phe Phe Lys Ile Asn Thr $195 \hspace{1.5cm} 200 \hspace{1.5cm} 205$

His Val Val Tyr Asn His Gln Glu Gln Ala Lys Tyr Glu Asn His Thr 210 215 220

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ccc Pro	aaa Lys 20	tgt Cys	gat Asp	tta Leu	aca Thr	aaa Lys 25	gaa Glu	aat Asn	gca Ala	ctt Leu	tcc ser 30	caa Gln	ata Ile	aca Thr	aac Asn	152
cta Leu 35	caa Gln	aca Thr	ccc Pro	aca Thr	aac Asn 40	aaa Lys	tta Leu	ttc Phe	atc Ile	aaa Lys 45	att Ile	tgc Cys	aga Arg	gaa Glu	cta Leu 50	200
cat His	gaa Glu	aat Asn	ggg Gly	gaa Glu 55	Pro	cat His	ctc Leu	cat His	att Ile 60	ttg Leu	atc Ile	caa Gln	ttc Phe	gaa Glu 65	gga Gly	248
aaa Lys	tac Tyr	aat Asn	tgt Cys 70	acc Thr	aat Asn	caa Gln	cga Arg	ttc Phe 75	ttc Phe	gac Asp	ctg Leu	gta Val	tcc Ser 80	cca Pro	acc Thr	296
agg Arg	tca Ser	gca Ala 85	cat His	ttc Phe	cat His	ccg Pro	aac Asn 90	att Ile	cag Gln	gga Gly	gct Ala	aaa Lys 95	tcg Ser	agc Ser	tcc Ser	344
Asp	gtc Val 100	aag Lys	tcc Ser	tat Tyr	atc Ile	gac Asp 105	aag Lys	gac Asp	gga Gly	gat Asp	gtt Val 110	ctt Leu	gaa Glu	tgg Trp	ggt Gly	392
act Thr 115	ttc Phe	cag Gln	atc Ile	Asp	gga Gly 120	cga Arg	tct Ser	gct Ala	agg Arg	gga Gly 125	gga Gly	caa Gln	cag Gln	aca Thr	gcc Ala 130	440

Pagina 8

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Thr Asn Leu Gln Thr Pro Thr Asn Lys Leu Phe Ile Lys Ile Cys Arg 40 45

Glu Leu His Glu Asn Gly Glu Pro His Leu His Ile Leu Ile Gln Phe 50 60

Glu Gly Lys Tyr Asn Cys Thr Asn Gln Arg Phe Phe Asp Leu Val Ser 65 75 80

Pro Thr Arg Ser Ala His Phe His Pro Asn Ile Gln Gly Ala Lys Ser 85 90 95

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Pagina 9

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tggatg	gatg	aaaacataaa	aaaacaaaat	catactaacc	aagtgatgtt	tttccttgtt	420
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INTERNATIONAL SEARCH REPORT

Inter inal Application No PC . , T2004/000287

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A. CLASSI IPC 7	FICATION OF SUBJECT MATTER C12N15/82 C07K14/01		
According to	o International Patent Classification (IPC) or to both national classifica	ation and IPC	
B. FIELDS	SEARCHED		
Minimum do IPC 7	ocumentation searched (classification system followed by classification ${\tt C12N-C07K}$	on symbols)	
Documentat	ion searched other than minimum documentation to the extent that s	uch documents are included in the fields se	arched
Electronic d	ata base consulted during the international search (name of data base	se and, where practical, search terms used)	
EPO-In	ternal, BIOSIS, EMBASE, Sequence Sea	rch, PAJ, WPI Data	
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the rek	evant passages	Relevant to claim No.
A	WO 00/43520 A (ABOUZID AHMED M; ERNEST (US); HUNTER WAYNE B (US); JA) 27 July 2000 (2000-07-27) the whole document		1-43
Α	BRUNETTI ANGELA ET AL: "Transgen expressed T-Rep of tomato yellow sardinia virus acts as a trans-dominant-negative mutant, i viral transcription and replicati JOURNAL OF VIROLOGY, vol. 75, no. 22, November 2001 (2 pages 10573-10581, XP002300782 ISSN: 0022-538X the whole document	leaf curl nhibiting on"	1-43
X Furth	ner documents are listed in the continuation of box C.	Patent family members are listed in	n annex.
"A" docume consid "E" earlier of filing d "L" docume which citation "O" docume other r	ent defining the general state of the art which is not ered to be of particular relevance document but published on or after the international ate nt which may throw doubts on priority claim(s) or is cited to establish the publication date of another or other special reason (as specified) ent referring to an oral disclosure, use, exhibition or neans ent published prior to the international filing date but	 "T" later document published after the Inter or priority date and not in conflict with the cited to understand the principle or the invention. "X" document of particular relevance; the clannot be considered novel or cannot involve an inventive step when the document of particular relevance; the clannot be considered to involve an involve an involve an involve an involve an involve and involve an involve and involve an involve and involve an	the application but ory underlying the alimed Invention be considered to current is taken alone alimed Invention entive step when the re other such docu- s to a person skilled
	actual completion of the international search	Date of mailing of the international sear	ch report
-	5 October 2004	02/11/2004	
Name and n	nalling address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL – 2280 HV Rijswijk	Authorized officer	
	Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Mundel, C	

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inte nal Application No
P(___T2004/000287

		P(T2004/000287
C.(Continua	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
А	NORIS E ET AL: "Resistance to tomato yellow leaf curl geminivirus in Nicotiana benthamiana plants transformed with a truncated viral C1 gene" VIROLOGY, ACADEMIC PRESS,ORLANDO, US, vol. 224, no. 1, 1996, pages 130-138, XP002142359 ISSN: 0042-6822 the whole document	1-43
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	Patent document cited in search report		Publication date		Patent family member(s)		Publication date	
	WO 0043520	Α	27-07-2000	AU WO	285640 004352	00 A 20 A2	07-08-2000 27 - 07-2000	
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Form PCT/ISA/210 (patent family ennex) (January 2004)